# Sensitivity Analysis

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#### ABSTRACT

This study investigates the process of caraway oil extraction from chamomile flowers. A parameter-distributed model consisting of a set of partial differential equations is used to describe the governing mass transfer phenomena in a solid-fluid environment under supercritical conditions using carbon dioxide as a solvent. The concept of quasi-one-dimensional flow is applied to reduce the number of spatial dimensions. The flow is assumed to be uniform across any cross-section, although the area available for the fluid phase can vary along the extractor. The physical properties of the solvent are estimated from the Peng-Robinson equation of state. A set of laboratory experiments was performed under multiple constant operating conditions:  $30-40^{\circ}C$ , 100-200 bar, and  $3.33-6.67\cdot10^{-5}$  kg/s. Different sensitivity analysis methods can be applied to assess the robustness of the model parameters and their influence on the process model. The local sensitivity analysis investigates the influence of infinitely small changes in the model parameters and the controls on model output. This work focuses on analysing the effect of pressure on the model state space and the extraction yield.

## 1. Introduction

This study investigates the extraction of essential oil from chamomile flowers (Matricaria chamomilla L.) via supercritical fluid extraction techniques and the modelling of this process. Chamomile is a medicinal herb widely cultivated in southern and eastern Europe — in countries such as Germany, Hungary, France and Russia. It can be found outside Europe, for instance in Brazil as discussed by Singh et al. [1]. This plant is distinguished by its hollow, bright gold cones, housing disc or tubular florets and surrounded by about fifteen white ray or ligulate florets. Chamomile has been used for its medicinal benefits, serving as an anti-inflammatory, antioxidant, mild astringent, and healing remedy. Aqueous extract of chamomile is widely used to calm nerves and mitigate anxiety, hysteria, nightmares, insomnia and other sleep-related conditions, according to Srivastava [2]. Orav et al. [3] reported that oil yields from dried chamomile samples ranged from 0.7 to 6.7 mL/kg. The highest yields of essential oil, between 6.1 and 6.7 mL/kg, were derived from chamomile sourced from Latvia and Ukraine. In comparison, chamomile from Armenia exhibited a lower oil content of 0.7 mL/kg.

Evaluating the economic viability of the process is essential when choosing a suitable technology for essential oil extraction. Traditional methods, such as distillation and organic solvent extraction, are commonly employed but have drawbacks. Distillation, for example, involves high temperatures that can lead to the thermal degradation of heat-sensitive compounds. This limitation has led to the increased popularity of alternative techniques such as supercritical fluid extraction. Supercritical carbon dioxide is appealing thanks to its distinctive properties: it is inflammable, non-toxic and non-corrosive. Supercritical fluids can exhibit

both gas- and liquid-like properties, allowing for adjustable dissolving power through changes in operating conditions.

The literature offers various mathematical models to describe the extraction of valuable compounds from biomass. Selecting a process model is case-to-case dependent and requires analysis of each model's specific assumptions about mass transfer and thermodynamic equilibrium.

The model proposed by Reverchon et al. [4] is called the hot ball model, as it is based on an analogy to heat transfer and describes an extraction process from solid particles. This model assumes that particles contain low quantities of solute and that solubility is not a limiting factor.

The Broken-and-Intact Cell model, proposed by Sovova [5], assumes that external surfaces of particles are mechanically disrupted, allowing the solvent access to the solute in the broken cells. In contrast, the solute in intact cells remains less accessible due to higher mass transfer resistance.

Reverchon [6] formulated a fluid-solid extraction model where the solute is treated as a single component, governed by internal mass transfer resistance and omitting the effects of external mass transfer, axial dispersion and variations in fluid density and flow rate throughout the bed.

This work builds upon the linear kinetic model suggested by Reverchon [6], deriving fundamental governing equations to develop a comprehensive model for the chamomile oil extraction process. This model aims for control-oriented simplicity, assuming a semi-continuous operation within a cylindrical vessel. The process involves a supercritical solvent being pumped through a fixed bed of finely chopped biomass to extract the solute, followed by separation of the solvent and solute in a flush drum to collect the extract. Parameters such as pressure (P), feed flow rate  $(F_{in})$  and inlet temperature  $(T_{in})$  are adjustable and measurable, while the outlet temperature  $(T_{out})$  and the amount of product at the outlet can only be monitored. Figure 1 presents a simplified process flow diagram.

This study aims to analyse the influence of changes in operating conditions on the process model described in

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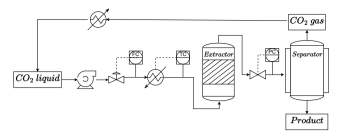


Figure 1: Process flow diagram

article 1. The relation between input and output is obtained by applying a sensitivity analysis, which examines the impact of model parameters or controls on the model output. The result of sensitivity analysis can be used to allocate the source of uncertainty in the system, perform the model reduction or search for errors in the model (by identifying unexpected relationships between inputs and outputs). There are many sensitivity analysis methods, which include but are not limited to those listed below:

- One-at-a-time method
- Derivative-based local methods
- Variance-based methods

In the literature, many different supercritical extraction models were analysed using different sensitivity analysis techniques. Fiori et al. [7], performed the sensitivity calculations by varying the parameters within their confidence interval and observing how the model results changed. Their sensitivity analysis revealed that changes in particle diameter and the internal mass transfer coefficient have the strongest influence on extraction in the second phase of the process (diffusion-control regime).

Santos et al. [8], in their work, considered the model of Sovova [5] for the semi-continuous isothermal and isobaric extraction processes using carbon dioxide as a solvent. The parametric sensitivity analysis was carried out using a factorial design in two levels. The model parameters were disturbed by 10%, and their main effects were analysed and strategies for high-performance operation are proposed. The authors performed the sensitivity analysis with respect to the superficial velocity, particle diameter, initial concentration of solute in the solid phase and the concentration of solute in the fluid phase at the inlet to the extractor.

Hatami and Ciftci [9] performed the one-factor-at-atime sensitivity analysis to assess the response of net present value concerning variations in both technical and economic variables. Their study is divided into two parts. In the first part, the sensitivity analysis is carried out for an extraction plant with a constant extractor volume of 300 L. The focus is on examining how net present value is influenced by changes in individual technical and economic parameters while keeping the extractor volume fixed. In the second part, the investigation revolves around the effects of altering the extractor volume (from 1 to 600 L) on the project's overall

profitability. Their findings show that the most influential factors on net present value include the price of the extract, the interest rate, the dynamic time, and the project lifetime.

#### 2. Materials and methods

# 2.1. Governing equations

Following the work of Anderson [11], the governing equations for a quasi-one-dimensional flow were derived. A quasi-one-dimensional flow refers to a fluid flow scenario assuming that the flow properties are uniformly distributed across any cross-section. This simplification is typically applied when the flow channel's cross-sectional area changes, such as through irregular shapes or partial filling of an extractor. According to this assumption, velocity and other flow properties change solely in the flow direction.

As discussed by Anderson [12], all flows are compressible, but some of them can be treated as incompressible when the Mach number is smaller than 0.3. This assumption leads to the incompressible condition:  $\nabla \cdot u = 0$ , which is valid for constant density (strict incompressible) or varying density flow. The assumption allows for removing acoustic waves and large perturbations in density and/or temperature. In the 1-D case, the incompressibility condition becomes  $\frac{du}{dz} = 0$ , so the fluid velocity is constant.

The set of quasi-one-dimensional governing equations in Cartesian coordinates is described by Equations 1 - 3:

$$\frac{\partial \left(\rho_f A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f v\right)}{\partial z} = 0 \tag{1}$$

$$\frac{\partial \left(\rho_f v A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f v^2\right)}{\partial z} = -A_f \frac{\partial P}{\partial z} \tag{2}$$

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$$\frac{\partial \left(\rho_f e A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f v e\right)}{\partial z} = -P \frac{\left(A_f v\right)}{\partial z} + \frac{\partial}{\partial z} \left(k \frac{\partial T}{\partial z}\right) \tag{3}$$

where  $\rho_f$  is the density of the fluid,  $A_f$  is the function which describes a change in the cross-section, v is the velocity, P is the total pressure, e is the internal energy of the fluid, t is time and z is the spatial direction.

## 2.2. Extraction model

## 2.2.1. Continuity equation

The previously derived quasi-one-dimensional continuity equation (Equation 1) is redefined by incorporating the function  $A_f = A\phi$ . This modification distinguishes constant and varying terms, where the varying term accounts for changes in the cross-sectional area available for the fluid. Equation 4 shows the modified continuity equation:

$$\frac{\partial(\rho_f\phi)}{\partial t} + \frac{\partial(\rho_f v A\phi)}{\partial z} = 0 \tag{4}$$

where A is the total cross-section of the extractor and  $\phi$ describes porosity along the extractor.

Assuming that the mass flow rate is constant in time, the temporal derivative becomes the mass flux F, and the spatial derivative can be integrated along z as

$$\int \frac{\partial (\rho_f v A \phi)}{\partial z} dz = F \to F = \rho_f v A \phi \tag{5}$$

To simplify the system dynamics, it is assumed that F is a control variable and affects the whole system instantaneously (due to  $\nabla \cdot u = 0$ ), which allows finding the velocity profile that satisfies mass continuity based on F,  $\phi$  and  $\rho_f$ :

$$v = \frac{F}{\rho_f A \phi} \tag{6}$$

Similarly, superficial velocity may be introduced:

$$u = v\phi = \frac{F}{\rho_f A} \tag{7}$$

The fluid density  $\rho_f$  can be obtained from the Peng-Robinson equation of state if the temperature and thermodynamic pressure are known along z. Variation in fluid density may occur due to pressure or inlet temperature changes. In a non-isothermal case, in Equations 6 and 7  $\rho_f$  is considered the average fluid density along the extraction column.

## 2.2.2. Mass balance for the fluid phase

Equation 8 describes the movement of the solute in the system, which is constrained to the axial direction due to the quasi-one-dimensional assumption. Given that the solute concentration in the solvent is negligible, the fluid phase is described as pseudo-homogeneous, with properties identical to those of the solvent itself. It is also assumed that the thermodynamic pressure remains constant throughout the device. The analysis further simplifies the flow dynamics by disregarding the boundary layer near the extractor's inner wall. This leads to a uniform velocity profile across any cross-section perpendicular to the axial direction. Thus, the mass balance equation includes convection, diffusion and kinetic terms representing the fluid phase behaviour:

$$\frac{\partial c_f}{\partial t} + \frac{1}{\phi} \frac{\partial \left( c_f u \right)}{\partial z} = \frac{1 - \phi}{\phi} r_e + \frac{1}{\phi} \frac{\partial}{\partial z} \left( D_e^M \frac{\partial c_f}{\partial z} \right) \tag{8}$$

where  $c_f$  represents the solute concentration in the fluid phase,  $r_e$  is the mass transfer kinetic term and  $D_e^M$  is the axial diffusion coefficient.

## 2.2.3. Mass balance for the solid phase

As given by Equation 9, the solid phase is considered stationary, without convection and diffusion terms in the mass balance equation. Therefore, the only significant term in this equation is the kinetic term of Equation 10, which connects the solid and fluid phases. For simplicity, the extract is represented by a single pseudo-component:

$$\frac{\partial c_s}{\partial t} = \underbrace{r_e}_{\text{Kinetics}} \tag{9}$$

### 2.2.4. Kinetic term

As the solvent flows through the bed,  $\mathrm{CO}_2$  molecules diffuse into the pores and adsorb on the particle surface to form an external fluid film around the solid particles due to the solvent-solid matrix interactions. The dissolved solute diffuses from the particle's core through the solid-fluid interface, the pore and the film into the bulk. Figure 2 shows the mass transfer mechanism, where the mean solute concentration in the solid phase is denoted as  $c_s$ , and the equilibrium

concentrations at the solid-fluid interface are denoted as  $c_s^*$  and  $c_p^*$  for the solid and fluid phases, respectively. The concentration of the solutes in the fluid phase in the centre of the pore is denoted as  $c_p$ . As the solute diffuses through the pore, its concentration changes and reaches  $c_{pf}$  at the pore opening. Then, the solute diffuses through the film around the particle and reaches bulk concentration  $c_f$ . The two-film theory describes the solid-fluid interface inside the pore. The overall mass transfer coefficient can be determined from the relationship between the solute concentration in one phase and its equilibrium concentration.

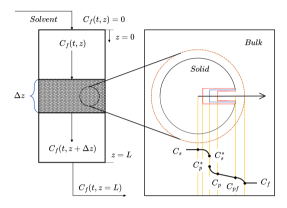


Figure 2: Mass transfer mechanism.

Bulley et al. [13] suggest a process where the driving force for extraction is given by the difference between the concentration of the solute in the bulk,  $c_f$ , and in the centre of the pore,  $c_p^*$ . The concentration  $c_p^*$  is in equilibrium with  $c_s$  according to the equilibrium relationship. The rate of extraction is thus  $r_e\left(c_f-c_p^*(c_s)\right)$ . In contrast, Reverchon [6] proposes a driving force given by the difference between  $c_s$  and  $c_p^*$ . Concentration  $c_p^*$  is determined by the equilibrium relationship with  $c_f$  and the extraction rate given by Equation 10:

$$r_e = \frac{D_i}{\mu l^2} \left( c_s - c_p^* \right) \tag{10}$$

where  $\mu$  is sphericity, l a characteristic dimension of particles that can be defined as l = r/3, r is the mean particle radius,  $\rho_s$  is the solid density,  $D_i$  corresponds to the overall diffusion coefficient and  $c_P^*$  is the concentration at the solid-fluid interface (which according to the internal resistance model is supposed to be at equilibrium with the fluid phase).

According to Bulley et al. [13], a linear equilibrium relationship (Equation 11) can be used to find the equilibrium concentration of the solute in the fluid phase  $c_f^*$  based on the concentration of the solute in the solid phase  $c_s$ :

$$c_f^* = k_p c_s \tag{11}$$

The volumetric partition coefficient  $k_p$  acts as an equilibrium constant between the solute concentration in one phase and the corresponding equilibrium concentration at the solid-fluid interphase. According to Spiro and Kandiah

[14],  $k_p$  can be expressed through the mass partition coefficient  $k_m$ :

$$k_m = \frac{k_p \rho_s}{\rho_f} \tag{12}$$

According to Reverchon [6], the kinetic term becomes

$$r_e = -\frac{D_i}{\mu l^2} \left( c_s - \frac{\rho_s c_f}{k_m \rho_f} \right) \tag{13}$$

## 2.2.5. Uneven solute's distribution in the solid phase

Following the idea of the Broken-and-Intact Cell (BIC) model (Sovova [15]), the internal diffusion coefficient  $D_i$  is considered to be a product of the reference value of  $D_i^R$  and the exponential decay function  $\gamma$ , as given by Equation 14:

$$D_i = D_i^R \gamma(c_s) = D_i^R \exp\left(\Upsilon\left(1 - \frac{c_s}{c_{s0}}\right)\right)$$
 (14)

where  $\Upsilon$  describes the curvature of the decay function. Equation 15 describes the final form of the kinetic term:

$$r_e = -\frac{D_i^R \gamma}{\mu l^2} \left( c_s - \frac{\rho_s c_f}{k_m \rho_f} \right) \tag{15}$$

The  $\gamma$  function limits the solute's availability in the solid phase. Similarly to the BIC model, the solute is assumed to be contained in the cells, some of which are open because the cell walls were broken by grinding, with the rest remaining intact. The diffusion of the solute from a particle's core takes more time than the diffusion of the solute close to the outer surface. The same idea can be represented by the decaying internal diffusion coefficient, where the decreasing term is a function of the solute concentration in the solid.

Alternatively, the decay function  $\gamma$  can be interpreted by referring to the Shrinking Core model presented by Goto et al. [16], where the particle radius changes as the amount of solute in the solid phase decreases. As the particle size decreases due to dissolution, the diffusion path increases, which makes the diffusion slower and reduces the value of the diffusion coefficient. This analogy can be applied to Equation 14 to justify the application of a varying diffusion coefficient.

# 2.2.6. Heat balance

The heat balance equation describe the evolution of the enthalpy in the system and it is given by Equation 16

$$\frac{\partial \left(\rho_f h A_f\right)}{\partial t} = -\frac{\partial \left(\rho_f h A_f v\right)}{\partial z} + \frac{\partial \left(P A_f\right)}{\partial t} + \frac{\partial}{\partial z} \left(k \frac{\partial T}{\partial z}\right) \tag{16}$$

The main advantage of this formulation is the presence of term  $\partial P/\partial t$ , which directly affects the system through the change of thermodynamic pressure.

If the value of enthalpy h is known from the time evolution of the energy equation, and pressure P is known from measurement, then the temperature T can be reconstructed based on the departure function. The departure function is a mathematical function that characterizes the deviation of a thermodynamic property (enthalpy, entropy, and internal energy) of a real substance from that of an ideal gas at the same temperature and pressure. As presented by Gmehling

et al. [17], for the Peng-Robinson equation of state, the enthalpy departure function is defined by Equation 17.

$$h - h^{id} = RT \left[ T_r(Z - 1) - 2.078(1 + \kappa) \sqrt{\alpha(T)} \ln \left( \frac{Z + \left( 1 + \sqrt{2} \right) B}{Z + \left( 1 - \sqrt{2} \right) B} \right) \right] (17)$$

Equation 17 requires an reference sate, which is assumed to be  $T_{ref} = 298.15$  K and  $P_{ref} = 1.01325$  bar.

A root-finder can be used to find a value of temperature, which minimizes the difference between the value of enthalpy coming from the heat balance and the departure functions. The root fining procedure to repeated at every time step to find a temperature profile along spatial direction z.

$$\min_{T} \left( \underbrace{h(t,x)}_{\text{Heat balance}} - \underbrace{h(T,P,\rho_f(T,P))}_{\text{Departure function}} \right)^2 \tag{18}$$

## 2.2.7. Pressure term

As explained in Chapters 2.1, at Low-Mach number conditions, the thermodynamic pressure is nearly constant in space due to the small pressure wave propagation that occurs at the speed of sound. Under such conditions, the term  $\partial P/\partial t$  can be approximated by a difference equation, which describes the pressure change in the whole system. The pressure P in the system is considered a state variable, while the pressure in the new time-step  $P_{in}$  is considered a control variable.

$$\frac{\partial P}{\partial t} \approx \frac{P_{in} - P}{\Delta t} \tag{19}$$

Such a simplified equation allows for instantaneous pressure change in the system but does not consider a gradual pressure build-up and the effects of pressure losses. In a real system, the dynamics of pressure change would depend on a pump and a back-pressure regulator.

# 2.2.8. Extraction yield

The process yield is calculated according to Equation 20 as presented by Sovova et al. [18]. The measurement equation evaluates the solute's mass at the extraction unit outlet and sums it up. The integral form of the measurement (Equation 20) can be transformed into the differential form (Equation 21) and augmented with the process model.

$$y = \int_{t_0}^{t_f} \frac{F}{\rho_f} c_f \Big|_{z=L} dt \tag{20}$$

$$\frac{dy}{dt} = \left. \frac{F}{\rho_f} c_f \right|_{z=L} \tag{21}$$

#### 2.2.9. Initial and boundary conditions

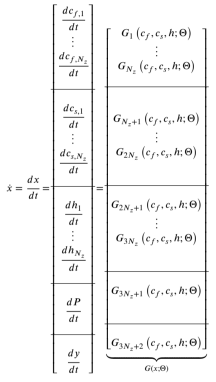
It is assumed that the solvent is free of solute at the beginning of the process  $c_{f0} = 0$ , that all the solid particles have the same initial solute content  $c_{s0}$ , and that the system is isothermal, hence the initial state is  $h_0$ . The fluid at the inlet is considered not to contain any solute. The initial and boundary conditions are defined as follows:

$$c_f(t=0,z) = 0 \qquad c_s(t=0,z) = c_{s0} \qquad h(t=0,z) = h_0$$

$$\begin{split} c_f(t,z=0) &= 0 \qquad h(t,z=0) = h_{in} \qquad \frac{\partial c_f(t,z=L)}{\partial x} = 0 \\ \frac{\partial h(t,z=L)}{\partial x} &= 0 \qquad c_s(t,z=\{0,L\}) = 0 \qquad y(0) = 0 \quad P(0) = P_0 \end{split}$$

# 2.2.10. Discretization methods

The method of lines is used to transform the process model equations into a set of ODEs denoted as  $G(x;\Theta)$ . The backward finite difference is used to approximate the first-order derivative, while the central difference scheme approximates the second-order derivative z direction. The length of the fixed bed is divided into  $N_z$ , i.e. equally distributed points in the z direction. The state-space model after discretization is denoted as x and defined as follows:



where  $x \in \mathbb{R}^{N_x=3N_z+2}$  and  $\Theta \in \mathbb{R}^{N_\Theta=N_\theta+N_u}$ ,  $N_\theta$  is the number of parameters,  $N_u$  is the number of control variables.

For a derivative to be conservative, it must form a telescoping series. In other words, only the boundary terms should remain after adding all terms coming from the discretization over a grid, and the artificial interior points should be cancelled out. Discretization is applied to the conservative form of the model to ensure mass conservation.

# 2.3. Local sensitivity analysis

Local derivative-based methods involve taking the total derivative of the state vector x with respect to the parameter space  $\Theta$ . The set of derivatives, known as sensitivity equations, is integrated simultaneously with the process model. The sensitivity analysis shows how responsive the solution is for changes in the parameter  $\Theta$ . As discussed by Dickinson and Gelinas [19], the sensitivity analysis can be used to determine the influence of the uncertainty on the solution of the original system. A sensitivity analysis can be used

to distinguish sensitive parameters from insensitive ones, which might be helpful for model reduction. Finally, from a control engineering point of view, the sensitivity analysis allows sorting the control variables with respect to the level of effort required to change the model's output.

Following the work of Maly and Petzold [20], the sensitivity equations  $(\dot{S})$  can be defined as follow:

$$S(x;\Theta) = \frac{dx}{d\Theta} \tag{22}$$

The new system of equations can be obtained by taking derivatives of S with respect to time t and applying the chain rule.

$$\dot{S}(x;\Theta) = \frac{dS(x;\Theta)}{dt} = \frac{d}{dt} \left(\frac{dx}{d\Theta}\right) = \frac{d}{d\Theta} \left(\frac{dx}{dt}\right) = \frac{dG(x;\Theta)}{d\Theta}$$
(23)

The sensitivity Equation 24 can be obtained by applying the definition of the total derivative to Equation 23.

$$\frac{dG(x;\Theta)}{d\Theta} = \underbrace{\frac{\partial G(x;\Theta)}{\partial x}}_{J_x(x;\Theta)} \underbrace{\frac{\partial x}{\partial \Theta}}_{S(x;\Theta)} + \underbrace{\frac{\partial G(x;\Theta)}{\partial \Theta}}_{J_{\Theta}(x;\Theta)}$$
(24)

The Jacobian  $J_x(x;\Theta)$  represents the matrix of equations of size  $N_x \times N_x$ , where each equation  $J_x(n_x,n_x)$  is the derivative of  $G_{n_x}(x;\Theta)$  with respect to the state variable  $x_{n_{\Theta}}$ .

$$J_{x}(x;\Theta) = \begin{pmatrix} \frac{\partial G_{1}(x;\Theta)}{\partial x_{1}} & \frac{\partial G_{1}(x;\Theta)}{\partial x_{2}} & \dots & \frac{\partial G_{1}(x;\Theta)}{\partial x_{N_{x}}} \\ \frac{\partial G_{2}(x;\Theta)}{\partial x_{1}} & \frac{\partial G_{2}(x;\Theta)}{\partial x_{2}} & \dots & \frac{\partial G_{2}(x;\Theta)}{\partial x_{N_{x}}} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial G_{N_{x}}(x;\Theta)}{\partial x_{1}} & \frac{\partial G_{N_{x}}(x;\Theta)}{\partial x_{2}} & \dots & \frac{\partial G_{N_{x}}(x;\Theta)}{\partial x_{N_{x}}} \end{pmatrix}$$
(25)

The sensitivity matrix  $S(x;\Theta)$  represents the matrix of equations of size  $N_x \times N_\Theta$ , where each subequation  $S(n_x,n_\Theta)$  is the derivative of the state variable  $x_{n_x}$  with respect to the parameter  $\Theta_{n_\Theta}$ . Matrix  $J_x(x;\Theta)$  and  $S(x;\Theta)$  describe indirect influence of  $\Theta_{n_\Theta}$  on the state space.

$$S(x;\Theta) = \begin{pmatrix} \frac{\partial x_1}{\partial \Theta_1} & \frac{\partial x_1}{\partial \Theta_2} & \cdots & \frac{dx_1}{\partial \Theta_{N_{\Theta}}} \\ \frac{\partial x_2}{\partial \Theta_1} & \frac{\partial x_2}{\partial \Theta_2} & \cdots & \frac{dx_2}{\partial \Theta_{N_{\Theta}}} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial x_{N_x}}{\partial \Theta_1} & \frac{\partial x_{N_x}}{\partial \Theta_2} & \cdots & \frac{\partial x_{N_x}}{\partial \Theta_{N_{\Theta}}} \end{pmatrix}$$

$$(26)$$

The Jacobian  $J_{\Theta}(x;\Theta)$  represents the matrix of equations of size  $N_x \times N_{\Theta}$ , where each subequation  $J_{\Theta}(n_x,n_{\Theta})$  is the partial derivative of the process model equation  $G_{n_x}$  with respect to the parameter  $\Theta_{n_{\Theta}}$ .  $J_{\Theta}(n_x,n_{\Theta})$  defines direct effect of  $\Theta_{n_{\Theta}}$  on the state space.

$$J_{\Theta}(x;\Theta) = \begin{pmatrix} \frac{\partial G_{1}(x;\Theta)}{\partial \Theta_{1}} & \frac{\partial G_{1}(x;\Theta)}{\partial \Theta_{2}} & \dots & \frac{\partial G_{1}(x;\Theta)}{\partial \Theta_{N_{\Theta}}} \\ \frac{\partial G_{2}(x;\Theta)}{\partial \Theta_{1}} & \frac{\partial G_{2}(x;\Theta)}{\partial \Theta_{2}} & \dots & \frac{\partial G_{2}(x;\Theta)}{\partial \Theta_{N_{\Theta}}} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial G_{N_{x}}(x;\Theta)}{\partial \Theta_{1}} & \frac{\partial G_{N_{x}}(x;\Theta)}{\partial \Theta_{2}} & \dots & \frac{\partial G_{N_{x}}(x;\Theta)}{\partial \Theta_{N_{\Theta}}} \end{pmatrix}$$
(27)

The augmented system containing the original set of equations  $G(x; \Theta)$  and sensitivity equations can be formulated as  $G(x; \Theta)$ . The size of  $G(x; \Theta)$  is equal to  $N_s = N_x(N_{\Theta} + 1)$ .

$$\mathbf{G}\left(x;\Theta\right) = \begin{bmatrix} G(x;\Theta) \\ J_{x}(x;\Theta)S(x;\Theta) + J_{\Theta}(x;\Theta) \end{bmatrix} \tag{28}$$

The initial conditions are described as

$$\mathbf{G}\left(x(t_0);\Theta\right) = \begin{bmatrix} x(t_0), & \frac{\mathrm{d}x(t_0)}{d\Theta_1}, & \cdots, & \frac{dx(t_0)}{d\Theta_{N_{\Theta}}} \end{bmatrix}^{\mathsf{T}} = (29)$$

$$= \begin{bmatrix} x_0, & 0, & \cdots, & 0 \end{bmatrix}^{\mathsf{T}}$$
(30)

The sensitivity analysis of the output function can be performed with respect to parameters  $\Theta$ . The output function g(x) returns y. By taking a total derivative of y with respect to  $\Theta$ , the new sensitivity equation can be found.

$$\frac{dy}{d\Theta} = \frac{dg(x)}{d\Theta} = \frac{\partial g(x)}{\partial x} \frac{\partial x}{\partial \Theta} + \frac{\partial g(x)}{\partial \Theta}$$
(31)

## 3. Results

This study examines the impact of pressure change on the state space and extraction yield through the local sensitivity analysis. The details of the process model and parameters are delivered in article 1. The process model was validated under the following range of operating conditions: temperatures between  $30 - 40^{\circ}C$ , pressures between 100 - 200 bar, and mass flow rates between  $3.33 - 6.67 \cdot 10^{-5}$  kg/s. The system is assumed to operate at the midpoint of the validated range:  $35^{\circ}C$ , 150 bar and  $5 \cdot 10^{-5}$  kg/s.

A step-function shown in Figure 3 illustrates the pressure change in the extractor. As discussed in Chapter 2.1, a small pressure wave propagates at the speed of sound relative to the flow. If the flow velocity is relatively low, all pressure changes are hydrodynamic (resulting from velocity motion) rather than thermodynamic. The Low Mach number assumption leads to instant propagation of the thermodynamic pressure throughout the system and allows for considering a single pressure value for the entire system. Analogously to Figure 3, the pressure change affects the enthalpy balance by introducing a step-change along the whole system as presented in Figure 4.

The pressure change affects the fluid's temperature inside the computational domain, but boundary values are subject to constraints specified at the extremes of that domain. Applying the Dirichlet boundary conditions imposes a fixed temperature value at the inlet, which leads to a thermal gradient propagating along the system. In contrast, Neumann boundary conditions dictate the heat flux at the boundaries. In this work, Neumann boundary conditions of zero were used to ensure that the temperatures at the inlet, outlet, and inside the extractor would vary uniformly and according to the pressure change.

Figure 5 shows the sensitivity of the solute concentration in the solid phase in response to pressure change. As discussed in Chapter 2.2.1, the velocity of a fluid is inversely proportional to its density, which suggests that the pressure reduces the fluid's velocity. As a result, the residence

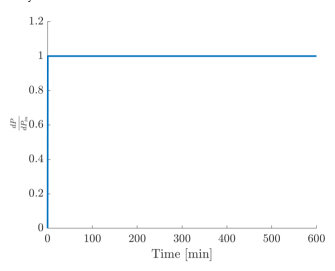
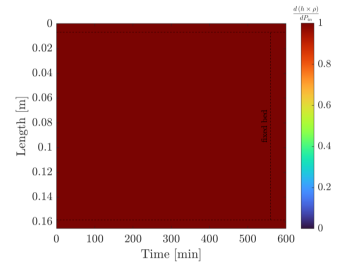


Figure 3: The effect of  $P_{in}$  change on P



**Figure 4**: The effect of  $P_{in}$  change on  $(h \times \rho)$ 

time is extended, causing longer interaction between the solute and solvent. Initially, the extraction process is in the kinetic-controlled regime, where the concentration gradient is high, and the limiting factor is the solute solubility. As discussed in article 1, the system is considered to be far from saturation, which explains the low system response at the beginning of the process. Fiori et al. [7] also observed low sensitivity at the beginning of extraction. The system response becomes evident when the concentration gradient diminishes, and the extraction switches from the kineticcontrolled to the diffusion-controlled regime. The negative sign can be interpreted as a faster solute loss from the solid phase, which corresponds to enhanced mass transfer. Over time, the amount of solute becomes a limiting factor, and the pressure change effect on the system is reduced. Eventually, the sensitivities approach zero asymptotically when the solute is washed out from the solid bed.

Figure 6 shows how sensitive is the solute concentration in the fluid phase to the pressure change. Compared to Figure

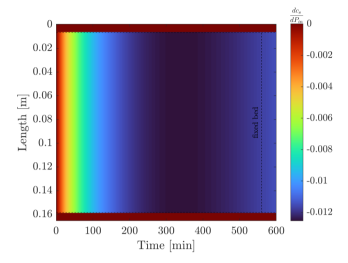


Figure 5: The effect of  $P_{in}$  change on  $C_s$ 

5, the dynamic behaviour of the fluid phase sensitivities can be observed. Due to the advection, the sensitivities move along the system analogously to the solute in the fluid phase. Although the pressure increase improves the mass transfer, the system response is initially low, which reflects the idle period discussed above. Later, the sensitivities increase due to faster solute loss from the solid phase. The positive sensitivities indicate that more solute is transported from the particles to the fluid. When the amount of solute in the solid phase becomes a limiting factor, the extraction rate slows down, and sensitivities decline asymptotically towards zero.

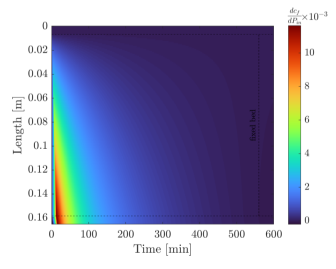


Figure 6: The effect of  $P_{in}$  change on  $C_f$ 

Figure 7 illustrates how sensitive the extraction yield is to the pressure change. Initially, the sensitivity curve stays almost flat, suggesting a latency in the system's response. Due to the decreased velocity of the fluid, the solute reaches the extractor's outlet later, which causes minor negative sensitivities to appear. When the solute reaches exits the

extractor, the sensitivity curve increases rapidly. The positive yield sensitivity indicates improved process efficiency, which is directly related to enhanced mass transfer. The peak in  $\frac{dy}{dP_{in}}$  reflects when the deviation from the original system is the largest. Eventually, the sensitivity declines and converges towards zero. The concentration gradient becomes a limiting factor, and the enhanced mass transfer no longer plays a dominant role compared to the original system.

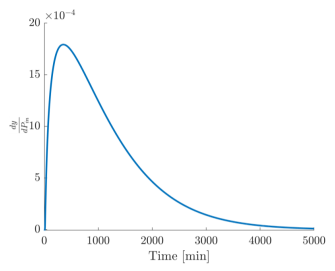


Figure 7: The effect of  $P_{in}$  change on y(t)

# 4. Conclusions

Sensitivity analysis is a tool to understand how model parameters affect its output. The presented formulation involves derivative-based local sensitivity analysis of the model solution with respect to selected parameters and controls. The local sensitivity analysis techniques consider only a small region of parameter space, and the conclusions derived from such an analysis are limited to local conditions. This work implemented the automatic differentiation to derive the sensitivity equations. In the case of dynamical systems, local sensitivity analysis provides a time series describing how that dependency evolves.

The local sensitivity analysis can be performed with respect to any model parameter, but this work focuses on the effect of pressure increase. At selected operating conditions  $(35^{\circ}C, 150 \text{ bar} \text{ and } 5 \cdot 10^{-5} \text{ kg/s})$ , the pressure increase enhances the mass transfer, which leads to faster loss of solute from particles and consequently to negative sensitivities in the solid phase. Analogously, the sensitivities in the fluid phase are characterised by positive deviations, which indicates that more solute is transported to the fluid phase. As a result, the extraction yield is improved and characterised by positive sensitivities. Local sensitivity analysis can provide valuable information by identifying which parameters are influential and how these influence changes over time. The sensitivity analysis results can be utilised for process modelling, experiment design, or model reduction.

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