## Parameter estimation

Oliwer Sliczniuk\*\*\*

<sup>a</sup>Aalto University, School of Chemical Engineering, Espoo, 02150, Finland

#### ARTICLE INFO

#### Keywords: Supercritical extraction Parameter estimation Mathematical modelling

#### ABSTRACT

Given a system of partial differential equations,  $F(t, x, \dot{x}, \Theta, u) = 0$ , where x represents state variables,  $\Theta$  are the parameters, and u are control variables, we describe the supercritical extraction process. The process model describes a partially filled extractor with a fixed bed, which work under constant operating conditions. We assume that the flow is uniform across any cross-section, although the area available for the fluid phase can change along the extractor. We apply the concept of quasi-one-dimensional flow to mimic the modelling of a two-dimensional case. In this work, a distributed-parameter model, based on Reverchon [1] is used to describe a fluid-solid extraction process of caraway oil from caraway seeds with  $CO_2$  as a solvent. The model parameters such as partition factor, internal diffusion coefficient, axial diffusion coefficient and saturation concentration are obtained from parameter estimation. The parameters are estimated based on four experiments performed at  $40^{\circ}C$  and  $50^{\circ}C$  at 200 bar and 300 bar. Given yield data, the model-based parameter estimation uses the maximum likelihood estimation method under assumption of normal error.

### 1. Introduction

The extraction of natural substances from solid materials and liquids with solvents have been a popular subject of research and development in the last years. Supercritical fluids have multiple applications in an extraction process due to the pressure-dependent dissolving power and both gas-and liquid-like properties (for example fluid-like density and gas-like diffusivity). Among different supercritical fluids, the supercritical  $CO_2$  is one of the most popular due to it is nontoxic, non-flammable and non-corrosive properties. The critical point of  $CO_2$  is relatively low (73.8 bar and 31°C), compare to other fluids. The supercritical extraction with  $CO_2$  become attractive alternative, to replace traditional extraction techniques.

The extraction of valuable compounds from fixed bed of biomass can be described by one of many mathematical models as presented by Huang et al. [2]. The selection the extraction model is not arbitrary and should be based on the knowledge of phenomenons occurring in the operational unit. Each model has its own assumptions and describe different mass transfer mechanisms and equilibrium relationships.

Based on analogy to heat transfer, Reverchon et al. [3] suggested a hot ball model, where the extraction process is treated analogously to a process in which a hot ball is cooled in a uniform medium. The hot ball model is used to describe an extraction process from solid particles which contains small quantities of solute so that the solubility is not a limiting factor.

Sovova [4] presented the Broken-and-Intact Cell model. The BIC model describes a system where the outer surfaces of particles have been mechanically interrupted. The solute from the broken cells is easily accessible for the solvent. The extraction of easily accessible solute is fast and directly

controlled by its diffusion and convection in the solvent. However, the rest of the solute is less accessible because it is closed in the particle's core or intact cells. This solute slowly diffuses through the walls of a cell due to high mass transfer resistance.

Reverchon [1] developed a model, which considers an oil as a single component and assumes that the extraction process is controlled by internal-mass transfer resistance. As a result of these assumptions, the external mass transfer was neglected. The original model of Reverchon does not consider the influence of axial dispersion and does not take into account the change of density and flow rate along the bed.

In our model, the extraction process is assumed to operate in a semi-continuous mode in a cylindrical vessel. The solvent is firstly brought to super-critical condition, pumped through a fixed-bed, of finely chopped biomass, where the solute is extracted from the biomass. Then, the solvent and solute from the extractor are separated in a flush drum, and the extract is collected. The flow rate  $(F_{in})$  and inlet temperature  $(T_{in})$  of the extractor's feed can be measured and manipulated. The pressure (P) in the vessel can be measured and manipulated, while the outlet temperature  $(T_{out})$  can only be measured. A simplified flow diagram is depicted in Figure 1.

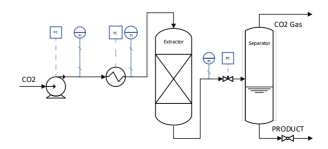


Figure 1: Process flow diagram (Font size)

<sup>\*</sup>Corresponding author

oliwer.sliczniuk@aalto.fi (O. Sliczniuk\*)
ORCID(s): 0000-0003-2593-5956 (O. Sliczniuk\*)

## 2. Materials and methods

## 2.1. Supercritical fluids

A supercritical fluid (SCF) is any substance at a temperature and pressure above its critical point, where distinct liquid and gas phases do not exist, but below the pressure required to compress it into a solid. It can effuse through porous solids like a gas, overcoming the mass transfer limitations that slow liquid transport through such materials. SCF are much superior to gases in their ability to dissolve materials like liquids or solids. Also, near the critical point, small changes in pressure or temperature result in large changes in density, allowing many properties of a supercritical fluid to be "fine-tuned". By changing the pressure and temperature of the fluid, the properties can be "tuned" to be more liquid-like or more gas-like.

The properties of a fluid can be divided into two kinds, equilibrium properties and transport properties. The equation of state can be used accurately to predict the equilibrium properties, such as density, enthalpy, vapor pressure, fugacity and fugacity coefficient, vapor liquid equilibrium, and all kinds of excess properties.

The thermodynamic properties of supercritical  $CO_2$  such as density, local speed of sound and specific heat capacity vary significantly for slight change in temperature and pressure due to real gas effects. Equation of state (EOS) which account these real gas effects is used to calculate the thermodynamic properties. In order to predict this real gas effects, the Peng-Robinson equation of state (P-R EOS) is used to compute properties of  $CO_2$ . Detail information about Peng-Robinson equation of state can be found in the work of Peng and Robinson [5], Elliott [6] or Pratt [7]. The P-R EOS belongs to the specific class of thermodynamic models for modelling the pressure of a gas as a function of temperature and density and can be written as a cubic function of the molar volume (of the density). The P-R EOS is presented by equation 1

$$P = \frac{RT}{V_m - b} - \frac{a\alpha}{V_m^2 + 2bV_m - b^2} \tag{1}$$

where  $a, b, \alpha$  are parameters defined as presented in the appendix A.1.1.

The properties of the  $CO_2$  presented as a function of operating conditions (temperature and pressure) are presented on Figure 2

At standard atmospheric pressure and temperature, the  $CO_2$  behaves as an ideal gas, and its compressibility factor equals to unity. However, at high temperature and pressure, the compressibility factor varies from unity, due to real gas effects. As it is presented on Figure 2a, the compressibility factor obtained from the Peng-Robinson equation of state varies strongly depending on the operating conditions. The compressibility factor can be obtained by given temperature and pressure by solving the polynomial form of the P-R EOS given by equation 2.

$$Z^{3} - (1 - B)Z^{2} + (A - 2B - 3B^{2})Z - (AB - B^{2} - B^{3}) = 0$$
 (2)

where A and B are parameters as defined in the appendix A.1.1. The roots of the polynomial can be found

iteratively or by the Cardano formula. One or three real roots can be found depending on the operating conditions. In a one-phase region, the fluid is described by one real root corresponding to the gas, liquid or supercritical phase. In the two-phase region, the gas-liquid mixture is present and has two roots. The biggest root is assigned to the gas phase, and the smallest root corresponds to the liquid phase.

The real gas effects are also visible on the density plot presented on the Figure 2b. The density calculations are based on the compressibility factor, as explained in the appendix A.1.2, and its value depends on the operating conditions.

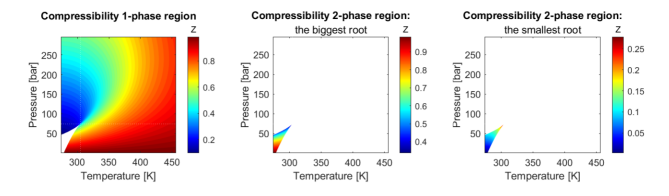
By analysing the compressibility and density plots more it can be noticed that very near to the critical point, the fluid can neither be called a liquid nor a gas and has a unique combination of gas-like and liquid-like properties.

The Figure 2c show behaviour of the heat capacity of a supercritical fluid at constant pressure  $(C_p)$ . The details of the calculations can be found in the appendix A.1.3. In contrary to the density which varies monotonically, the specific heat shows very high levels in a narrow region. In the subcritical region, the phase transition is associated with an effective spike in the heat capacity (i.e., the latent heat). Approaching the critical point, the latent heat falls to zero but this is accompanied by a gradual rise in heat capacity in the pure phases near phase transition. At the critical point, the latent heat is zero but the heat capacity shows a diverging singularity. Beyond the critical point, there is no divergence, but rather a smooth peak in the heat capacity; the highest point of this peak identifies the Widom line (as discussed by Simeoni et al. [8] and Banuti [9]).

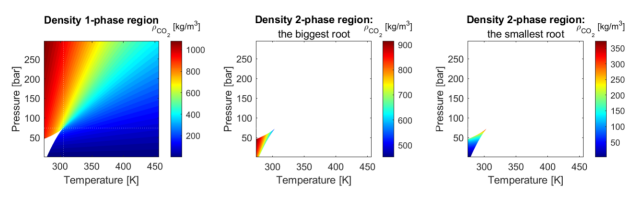
In order to calculate thermodynamic properties from a real gas, the departure function for that property with respect to the chosen equation of state has to be evaluated. As presented by Elliott [6], the departure function for any thermodynamic property of a real gas is defined as the difference in the value of that property determined from the chosen real gas equation of state and the value of the same property for an ideal gas under the same conditions of temperature and pressure.

On the other hand, transport properties (viscosity and conductivity) are also important quantities required in engineering design for production, fluid transportation, and processing. According to Sheng et al. [10], there is no satisfactory theory of transport properties of real dense gases and liquids. The main difficulties in the study of transport properties are twofold: one is the inherent difficulties involved in accurate measurements, and the other is the complexity involved in theoretical treatments. Therefore, the generally used correlations of transport coefficient are either empirical or based on some theoretical foundation. Enskog () developed a popular theory for the transport properties of dense gas based on the distribution function. However, the Enskog theory was proposed for rigid spherical molecules. For real gases, some modification is needed. Following the Enskog theory, many correlations have been proposed in the form of the reduced density and reduced temperature. The

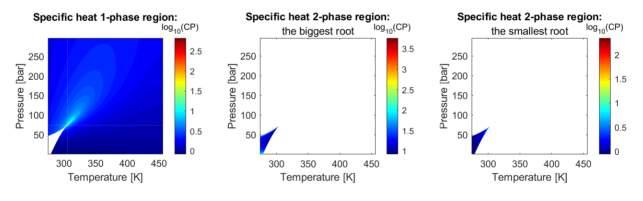
Add ref



(a) The compressibility factor based on the Peng-Robinson equation of state



(b) The fluid density based on the Peng-Robinson equation of state



(c) The specific heat of the  $CO_2$  based on the Peng-Robinson equation of state

Figure 2: Properties of  $CO_2$  based on the equation of state

correlations of Fenghour et al. [11] and Laesecke and Muzny [12] implemented and compared as presented on Figure 3. The viscosity formulation proposed by NIST consists of four contributions: (i) for the limit of zero density, (ii) for the initial density dependence, (iii) for the residual viscosity, and (iv) for the singularity of the viscosity at the critical point. The NIST correlation covers temperatures from 100 to 2000 K for gaseous  $CO_2$  and from 220 to 700 K with pressures

along the melting line up to  $8000\ MPa$  for compressed and supercritical liquid states.

Similarly, several correlations for thermal conductivity of  $CO_2$  were compared on Figure 4. The presented figures are focused around critical point, where the singularity is present. Similarities between specific heat and the thermal conductivity can be observed. The NIST correlation (Huber et al. [13]) captures the singular behaviour of thermal

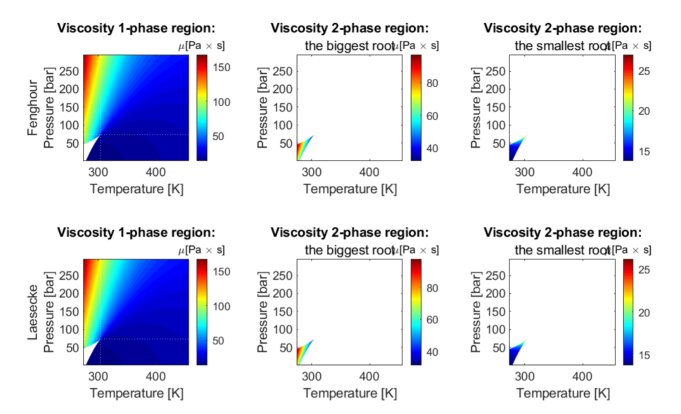


Figure 3: Viscosity obtained based on different correlations

conductivity around the critical point. The correlations are applicable for the temperature range from the triple point to  $1100\ K$  and pressures up to  $200\ MPa$ .

## 2.2. Governing equations

The detail derivation of the governing equation can be found in the appendix A.2 as well as in the work of Anderson [14]. Let's assume that any properties of the flow are unifrom across any given cross-section of an channel (or any device like an extractor). Such a flow is called quasi-one-dimensional. The variation of the cross-section might be an result of its irregular shape (or partial filling of an extractor).

The quasi-one-dimensional compressible Navier-Stokes equations in Cartesian coordinates is given be equations 3 to 5.

$$\frac{\partial \left(\rho_f A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f v\right)}{\partial z} = 0 \tag{3}$$

$$\frac{\partial \left( \rho_f v A_f \right)}{\partial t} + \frac{\partial \left( \rho_f A_f v^2 \right)}{\partial z} = -A_f \frac{\partial P}{\partial z} \tag{4}$$

$$\frac{\partial \left(\rho_f e A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f v e\right)}{\partial z} = -P \frac{\left(A_f v\right)}{\partial z} + \frac{\partial}{\partial z} \left(\frac{\partial T}{\partial z}\right) \tag{5}$$

where  $\rho_f$  is the density of the fluid,  $A_f$  is the function which describe change of 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 spacial direction.

Based on governing equations, the small discontinuity in flow properties, shown on figure 5, can be analysed. The analysis follow the work of Schreier [15].

The discontinuity is presumed to be at rest relative and the balance equations become

$$\begin{split} & \rho \delta v + v \delta \rho_f + \delta \rho_f \delta v = 0 \\ & \delta P = \delta v \delta \rho_f \end{split}$$

These relations are equally valid if the two regions are separated by a regions of finite width rather than a discontinuity.

$$\lim_{\rho_f v \to 0} \rho_f \delta v + v \delta \rho_f + \delta \rho_f \delta v = 0/\delta \rho_f \to \frac{dv}{d\rho_f} = -\frac{v}{\rho_f}$$

By combining momentum equation with above equation we get

$$\frac{dv}{d\rho_f} = -\frac{dv}{dP}\frac{dP}{d\rho_f} = -\frac{1}{\rho v}\frac{dP}{d\rho_f} = -\frac{v}{\rho_f}$$

If the flow is presumed to be isentropic,  $dP/d\rho = c^2$ , so  $V^2 = c^2$ , where c is the speed of sound. This can be interpreted that a small pressure wave propagates with the speed of sound relative to the flow.

## 2.3. Low Mach number expansion

As discussed by Lions [16], the low Mach number equations are a subset of the fully compressible equations of motion (continuity, momentum and energy). Such a

Fix the notation of pressure perturbation section to be coherent with governing equations

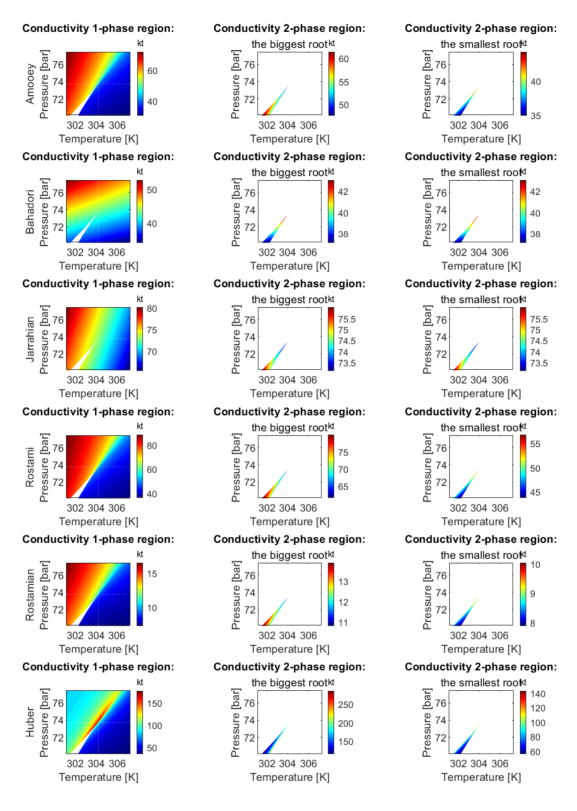


Figure 4: Thermal conductivity obtained based on different correlations

set of equations allow for large variations in gas density but it is consider to be acoustically incompressible. The low Mach number equations are preferred over the full compressible equations for low speed flow problems

 $\left(M_a = \frac{|V|}{\sqrt{\partial P/\partial \rho}} \ll 1\right)$  to avoids the need to resolve fast-moving acoustic signals. The equations are derived from the compressible equations based on the perturbation theory.

	ρ	$\rho + \delta \rho$	
$v \rightarrow$	P	$P + \delta P$	$v+\delta v\to$
	T	$T + \delta T$	

Figure 5: Small discontinuity in one-dimensional flow

The perturbation theory develops an expression for the desired solution in terms of a formal power series known as a perturbation series in some "small" parameter  $\zeta$ , that quantifies the deviation from the exactly solvable problem. The leading term in this power series is the solution of the exactly solvable problem, while further terms describe the deviation in the solution, due to the deviation from the initial problem.

The equations 3 to 4 describe the fully compressible equations of motion (respectively the transport of mass, momentum and energy) for quasi-one-dimensional case. We rescale the time variable, considering finally

$$\begin{split} \rho_{\zeta} &= \rho \left( z, t/M_a \right), \quad v_{\zeta} = \frac{1}{\zeta} v \left( z, t/M_a \right) \\ T_{\zeta} &= T \left( z, t/M_a \right), \quad k_{\zeta} = \zeta k(\rho_f, T) \end{split}$$

The conservative non-dimensional equations of motion becomes

$$\begin{split} &\frac{\partial \left(\rho_{\zeta}A_{f}\right)}{\partial t} + \frac{\partial \left(\rho_{\zeta}A_{f}v_{\zeta}\right)}{\partial z} = 0\\ &\frac{\partial \left(\rho_{\zeta}A_{f}v_{\zeta}\right)}{\partial t} + \frac{\partial \left(\rho_{\zeta}v_{\zeta}A_{f}v_{\zeta}\right)}{\partial z} + \frac{A_{f}}{M_{a}^{2}}\frac{\partial P_{\zeta}}{\partial z} = 0\\ &\frac{\partial \left(\rho_{\zeta}e_{\zeta}A_{f}\right)}{\partial t} + \frac{\partial \left(\rho_{\zeta}e_{\zeta}v_{\zeta}A_{f}\right)}{\partial z} - \frac{\partial}{\partial z}\left(k\frac{\partial T_{\zeta}}{\partial z}\right) + P_{\zeta}\frac{\partial \left(A_{f}v_{\zeta}\right)}{\partial z} = 0 \end{split}$$

Let's define  $\zeta=M_a^2$  and assume small Mach numbers,  $M_a\ll 1$ , then the kinetic energy, viscous work, and gravity work terms can be neglected in the energy equation since those terms are scaled by the square of the Mach number. The inverse of Mach number squared remains in the momentum equations, suggesting singular behaviour. In order to explore the singularity, the pressure, velocity and temperature are expanded as asymptotic series in terms of the parameter  $\zeta$ 

$$\begin{split} P_{\zeta} &= P_0 + P_1 \zeta + P_2 \zeta^2 + \mathcal{O}(\zeta^3) \\ \rho_{\zeta} &= \rho_0 + \rho_1 \zeta + \mathcal{O}(\zeta^2) \\ v_{\zeta} &= v_0 + v_1 \zeta + \mathcal{O}(\zeta^2) \\ T_{\zeta} &= T_0 + T_1 \zeta + \mathcal{O}(\zeta^2) \\ e_{\zeta} &= e_0 + e_1 \zeta + \mathcal{O}(\zeta^2) \end{split}$$

By expanding performing power expansion on the continuity equation and taking the limit of  $\zeta$  from the positive side we get

$$\begin{split} &\lim_{\zeta \to 0_{+}} \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \right)}{\partial t} + \\ &+ \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial z} = 0 \end{split}$$

The continuity equation become

$$\frac{\partial \left(\rho_0 A_f\right)}{\partial t} + \frac{\partial \left(\rho_0 A_f v_0\right)}{\partial z} = 0 \tag{6}$$

The form of the continuity equation stays the same. Considering the momentum equation, it can be seen that the inverse of Mach number squared remains which suggests singular behavior.

$$\begin{split} & \lim_{\zeta \to 0_{+}} \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial t} + \\ & + \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial z} + \\ & + A_{f} \frac{\partial}{\partial z} \left( \frac{P_{0}}{M_{a}^{2}} + \frac{P_{1} \zeta}{M_{a}^{2}} + \frac{P_{2} \zeta^{2}}{M_{a}^{2}} + \mathcal{O}(\zeta^{3}) \right) \end{split}$$

The first two terms stays the same, but the third one become different in structure. By further investigation of the pressure term in the momentum equation it can be observed

$$\begin{split} &\lim_{\zeta \to 0_+} \frac{\partial}{\partial z} \left( \frac{P_0}{M_a^2} + \frac{P_1 \zeta}{M_a^2} + \frac{P_2 \zeta^2}{M_a^2} + \mathcal{O}(\zeta^3) \right) = \\ &= \lim_{\zeta \to 0_+} \frac{\partial}{\partial z} \left( \frac{P_0}{M_a^2} \right) + \frac{\partial}{\partial z} \left( \frac{P_1 \zeta^2}{M_a^2} \right) + \frac{\partial}{\partial z} \left( \frac{P_2 \zeta^2}{M_a^2} \right) \\ &= \lim_{\zeta = M_a^2 \to 0_+} \frac{\partial}{\partial z} \left( \frac{P_0}{M_a^2} \right) + \frac{\partial}{\partial z} \left( \frac{P_1 M_a^2}{M_a^2} \right) + \frac{\partial}{\partial z} \left( \frac{P_2 M_a^4}{M_a^2} \right) \\ &= \lim_{\zeta = M_a^2 \to 0_+} 0 + \frac{\partial P_1}{\partial z} + 0 \end{split}$$

The simplification of the  $P_0$  in the momentum equation comes from the fact that  $P_0$  is independent of z. As it was presented above, the thermodynamical pressure moves with speed of sound, and any perturbation propagates instantaneously. The term related to  $P_2$  and higher order terms become zero at the limit of  $M_a \rightarrow 0$ . The momentum equation become

$$\frac{\partial \left(\rho_0 A_f v_0\right)}{\partial t} + \frac{\partial \left(\rho_0 v_0 A_f v_0\right)}{\partial z} + A_f \frac{\partial P_1}{\partial z} = 0$$

By expanding performing power expansion on the energy equation and taking the limit of  $\zeta$  from the positive side we get

$$\begin{split} &\lim_{\zeta \to 0_{+}} \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \left( e_{0} + e_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial t} + \\ &+ \frac{\partial \left( \left( \rho_{0} + \rho_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) A_{f} \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \left( e_{0} + e_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial z} + \\ &+ \frac{\partial}{\partial z} \left( k \frac{\partial}{\partial z} \left( T_{0} + T_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right) + \\ &- \left( P_{0} + P_{1} \zeta + P_{2} \zeta^{2} + \mathcal{O}(\zeta^{3}) \right) \frac{\partial \left( A_{f} \left( v_{0} + v_{1} \zeta + \mathcal{O}(\zeta^{2}) \right) \right)}{\partial z} = 0 \end{split}$$

The form of the energy equation stays the same.

$$\frac{\partial \left(\rho_{0}e_{0}A_{f}\right)}{\partial t}+\frac{\partial \left(\rho_{0}e_{0}v_{0}A_{f}\right)}{\partial z}-\frac{\partial}{\partial z}\left(k\frac{\partial T_{0}}{\partial z}\right)+P_{0}\frac{\partial \left(A_{f}v_{0}\right)}{\partial z}=0$$

where  $e_0 = e(\rho_0, T_0)$  and  $k = k(\rho_0, T_0)$ .

The expansion results in two different types of pressure and they are considered to be split into a thermodynamic

Add REF component  $(P_0)$  and a dynamic component  $(F_1)$ . The unermodynamic pressure is constant in space, but can change in time. The thermodynamic pressure is used in the equation of state. The dynamic pressure only arises as a gradient term in the momentum equation and acts to enforce continuity.

The resulting unscaled low Mach number equations are

$$\begin{split} &\frac{\partial \left(\rho_{f}A_{f}\right)}{\partial t} + \frac{\partial \left(\rho_{f}A_{f}v\right)}{\partial z} = 0 \\ &\frac{\partial \left(\rho_{f}A_{f}v\right)}{\partial t} + \frac{\partial \left(\rho_{f}vA_{f}v\right)}{\partial z} + A_{f}\frac{\partial P_{1}}{\partial z} = 0 \\ &\frac{\partial \left(\rho_{f}eA_{f}\right)}{\partial t} + \frac{\partial \left(\rho_{f}evA_{f}\right)}{\partial z} - \frac{\partial}{\partial z} \left(k\frac{\partial T}{\partial z}\right) + P_{0}\frac{\partial \left(A_{f}v\right)}{\partial z} = 0 \end{split}$$

The energy equation can be expanded through the chain rule to obtain

$$\rho A_{f}\left(\frac{\partial e}{\partial t} + v\frac{\partial e}{\partial z}\right) + e\underbrace{\left(\frac{\partial\left(\rho_{f}A_{f}\right)}{\partial t} + \frac{\partial\left(\rho_{f}vA_{f}\right)}{\partial z}\right)}_{Continutiy} - \frac{\partial}{\partial z}\left(k\frac{\partial T}{\partial z}\right) + P_{0}\frac{\partial\left(A_{f}v\right)}{\partial z} = 0$$
that

The non-conservative form of the energy equation become

$$\rho A_f \left( \frac{\partial e}{\partial t} + v \frac{\partial e}{\partial z} \right) - \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) + P_0 \frac{\partial \left( A_f v \right)}{\partial z} = 0$$

If the calorically perfect gas is assumed then  $e = C_n T$ , where  $C_n$  is the constant specific heat. The energy equation can derived in terms of temperature T

$$\rho A_f C_v \left( \frac{\partial T}{\partial t} + v \frac{\partial T}{\partial z} \right) - \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) + P_0 \frac{\partial \left( A_f v \right)}{\partial z} = 0$$

If isothermal case is assumed then, the energy equation becomes

$$\lim_{\Delta T \to 0_{+}} \rho A_{f} C_{v} \left( \frac{\partial T}{\partial t} + v \frac{\partial T}{\partial z} \right) - \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) + P_{0} \frac{\partial \left( A_{f} v \right)}{\partial z} = 0$$

which leads to

$$\frac{\partial \left(A_f v\right)}{\partial z} = 0\tag{7}$$

In one-dimensional case, the equation 7 become equivalent of  $\operatorname{div}(A_f v) = 0$ , which known as the incompressibility condition (Lions [16]).

A general formulation the internal energy for a real gas is:

$$de = C_v dT - \left[ P - T \left( \frac{\partial P}{\partial T} \right)_{v_m} \right] dv_m$$

where  $v_m$  is the molar volume.

The internal energy is a function of two intensive properties, in this case T and  $v_m = 1/\rho_f$ . But, in the case of an ideal gas, the equation of state is such that the second term in this equation is identically equal to zero. So the ideal gas is a special case in which the molar internal energy is a function only of temperature. For Peng-Robinson equation of state, the internal energy is defined as

$$e = C_v T + \frac{a\left(\alpha - T\frac{d\alpha}{dT}\right)}{2\sqrt{2}b} \ln \left[\frac{1 + b\left(1 - \sqrt{2}\rho\right)}{1 + b\left(1 + \sqrt{2}\rho\right)}\right]$$

Assuming constant temperature and pressure along space and in time:

$$\lim_{\Delta T, \Delta P \to 0_{+}} \rho A_{f} \left( \frac{\partial e}{\partial t} + \upsilon \frac{\partial e}{\partial z} \right) - \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) + P_{0} \frac{\partial \left( A_{f} \upsilon \right)}{\partial z} = 0 \rightarrow \frac{\partial \left( A_{f} \upsilon \right)}{\partial z} = 0$$

It can be deduce that the continuity equation becomes  $\frac{\partial \rho_f}{\partial t} = 0$  at constant temperature and pressure. Moreover incompressibility condition  $div(A_f V) = 0$  is obtained. = 0 at constant temperature and pressure. Moreover, the

Assuming an arbitrary function  $\hat{P}$ , which describe the total pressure, the dimensional momentum equation can be

$$V\left(\underbrace{\frac{\partial \left(\rho_{f} A_{f}\right)}{\partial t} + \frac{\left(\rho_{f} v A_{f}\right)}{\partial z}}_{Continuity equation}\right) + \rho_{f} A_{f} \frac{\partial v}{\partial t} + \rho_{f} v A_{f} \frac{\partial v}{\partial z} = -A_{f} \frac{\partial \hat{P}}{\partial z}$$

From the incompressibility conditions we can deduce

$$\frac{\partial (A_f v)}{\partial z} = 0 \to A_f \frac{\partial v}{\partial z} = -v \frac{\partial A_f}{\partial z}$$

By combining both above equations, assuming that

$$\frac{\rho_f v^2}{A_f} \frac{\partial A_f}{\partial z} = \frac{\partial \hat{P}}{\partial z} \to \int \frac{\rho_f v^2}{A_f} \frac{\partial A_f}{\partial z} dz = \int \frac{\partial \hat{P}}{\partial z} dz$$

The l.h.s integral can be solved by assuming  $\rho_f$  is constant and introducing superficial velocity  $u_s = A_f v$ 

$$\int \frac{\rho_f v^2}{A_f} \frac{\partial A_f}{\partial z} dz = \int \frac{\rho_f v^2}{A_f} \frac{A_f^2}{A_f^2} \frac{\partial A_f}{\partial z} dz$$

$$= \rho_f u_s^2 \int \frac{1}{A_f^3} \frac{\partial A_f}{\partial z} dz = -\frac{\rho_f u_s^2}{2\Delta A_f^2} = -\frac{\rho_f \Delta v^2}{2}$$

$$\int \frac{\partial \hat{P}}{\partial z} dz = \Delta \hat{P}$$

The final form of the momentum equation corresponds to the Bernoulli's principle

$$\Delta \hat{P} = -\frac{\rho_f \Delta v^2}{2} \xrightarrow{P_0 = \text{const}} \Delta M_a^2 P_1 = -\frac{\rho_f \Delta v^2}{2}$$

The Bernoulli's principle can be used to find the hydrodynamic pressure caused by varying cross-section at steadystate. Moreover, if the flow velocity is relatively low and so all pressure changes are hydrodynamic (due to velocity motion) rather than thermodynamic. The effect of this is that  $\partial \rho / \partial P = 0$ . In other words, the small changes in pressure due to flow velocity changes do not change the density. This has a secondary effect - the speed of sound in the fluid is  $\partial P/\partial \rho = \infty$  in this instance. So there is an infinite speed of sound, which makes the equations elliptic in nature. It can be deduced that at the isothermal conditions the density in the system propagates with the same speed as pressure due to the fact that they are both connected through the equation of state.

## 2.4. Extraction model

For the sake of clarity of the process model, different colors have been used in the equations to indicate: control variables, state variables, variables and parameters.

### 2.4.1. Continuity equation

The details of derivation can be found in the appendix A.2. The continuity equation for the fluid phase is given by equation 3. If  $A_f$  is specified to be a function of void fraction  $A_f = A\varepsilon(z)$ , where  $\varepsilon$  is void fraction of the bed and A is the cross-section of the empty extractor. The continuity equation become

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

Assuming that the mass flow-rate is constant in time, the temporal derivative become zero and the spacial derivative can be integrated along z.

$$\int \frac{\partial \left(\rho_f v \varepsilon\right)}{\partial z} dz = 0 \to F = \rho_f v \varepsilon \tag{8}$$

where F is a constant obtained from the integration and it is understood as the mass flux per unit area which is assumed to be constant along z. F can treated as a control variable, which can change in time but it is constant in space. Such a assumption allows to find the velocity profile which satisfy the mass continuity based on F,  $\varepsilon$  and  $\rho_f$ . The fluid density  $\rho_f$  density is connected through equation of state with thermodynamical pressure, which is assumed to be constant along z (due to the low-Mach number) and temperature. The variation in density might be caused by the accumulation of the fluid in the system (which happen instantaneously along z) or by temperature change. To simplify the dynamic of the system, it is assumed that F = F(t) is a control variable and affects the whole system instantaneously.

(a control variable) and temperature can be found from energy equation

Elaborate, the

pressure is know

Introduce relation between accumulation, pressure and energy equation

Double check if we can call it a plug flow

### 2.4.2. Mass balance for the fluid phase

The detail derivation of the mass balance equation for the fluid phase can be found in the appendix (A.2). The movement of the mobile pseudo-homogeneous phase (equation 9) is considered only in the axial direction. The properties of the system in the radial direction are assumed to be uniform. In addition, it is considered that the boundary layer adjacent to the inner wall of the extractor does not exist. Therefore, the velocity profile is constant across any cross-section of the extractor perpendicular to the axial direction. As a result, the plug flow model can be introduced. The particle size distribution and the void fraction of the solid phase cab change along an extractor but they remain constant in time. Moreover, the thermodynamic pressure is considered to be constant along the device, as discussed above. The amount of solute in the solvent is considered negligible. Therefore, the fluid phase can be described as pseudo-homogenous, and its properties are assumed to be the same as the solvent. The mass balance for the fluid phase consists of convection, diffusion, and kinetic terms.

$$\frac{\partial c_f(t,z)}{\partial t} = \underbrace{-\frac{u_s}{\varepsilon} \frac{\partial c_f(t,z)}{\partial z}}_{\text{Convective term}} - \underbrace{\frac{c_f(t,z)}{\varepsilon} \frac{\partial u(t,z)}{\partial z}}_{\text{Convective term}} + \underbrace{\frac{D_e^M}{\varepsilon} [T(t,z), P(t), F(t)]}_{\text{Diffusive term}} + \underbrace{\frac{\partial^2 c_f(t,z)}{\partial z^2}}_{\text{Kinetic term}} + \underbrace{\frac{1-\varepsilon}{\varepsilon} r_e(t,z)}_{\text{Kinetic term}}$$
(9)

where  $c_f(t,z), c_s(t,z), T(t,z)$  correspond to concentration of solute in the fluid phase, concentration of solute in the solid phase and the temperature, respectively.  $r_e(t,z)$  is a mass transfer kinetic term. F(t) is the mass flow rate, P(t) is the pressure, e is the void fraction of the bed, e0, e1, e2, e3, e4, e5, e6 is the solids density, e7, e8 is the solids density, e8, e9, e

Remove symbols which have been introduced before

## 2.4.3. Mass balance for the solid phase

Considering the solid phase to be fixed, the convection and diffusion terms in the corresponding mass balance (equation 10) are both assumed to be negligible or absent. Therefore, the only term present in this equation is the kinetic term (defined as presented in equation 11), which links solid and fluid phases. A single pseudo-component is used to represent the extract collectively.

$$\frac{\partial c_s(t,z)}{\partial t} = \underbrace{r_e(t,z)}_{\text{Kinetics}} \tag{10}$$

## 2.4.4. Kinetic term

The kinetic term is based on two-film theory and follow work of Reverchon [1]. The mass transfer kinetic (equation 11) consists of the overall diffusion coefficient and the concentration gradient, which acts as a driving force for the process.

As the solvent flows through the bed, the  $CO_2$  molecules diffuse into the pores and adsorb on the particle surface to for an external fluid film around the solid particles through the solvent-solid matrix interactions. Assuming that the mean free path of the molecule is much smaller than the pore diameter, the effect of Knudsen diffusion is small and can be neglected. The dissolved solute diffuses from the particle's core through the solid-fluid interface, the pore, and the film into the bulk. The graphical representation of the mass transfer mechanism is shown in Figure 6. The mean solute concentration in the solid phase is denoted as  $c_s$ . At the solid-fluid interface, the equilibrium concentrations are given as  $c_s^*$  and  $c_p^*$ , respectively for solid and fluid phases. 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  $c_{P,f}$ at the opening of the pore. The solute diffuses through the film around the particle and reaches a concentration in the bulk  $c_f$ . It can be assumed that the two-film theory describes the solid-fluid interface inside the pore. The overall mass transfer coefficient can be introduced if the relation between the solute concentration in one phase and its equilibrium concentration is known.

Bulley et al. [17] suggest a process where the driving force for extraction is given by the difference between 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 an equilibrium relationship. The rate of extraction is thus  $r_e$  ( $c_f - c_P^*(c_s)$ ).

On the other hand, Reverchon [1] proposes a driving force given by the difference between  $c_s$  and  $c_p^*$ .  $c_p^*$  is

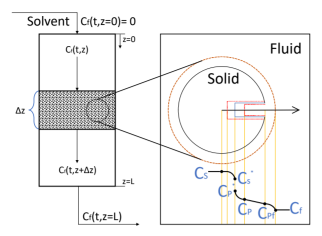


Figure 6: The extraction mechanism

determined by an equilibrium relationship with  $c_f$  and the extraction rate is  $r_e \left(c_s - c_p^*(c_f)\right)$  or more precisely

$$r_{e}(t,z) = \frac{D_{i}(T(t,z), P(t))}{\mu l^{2}} \left(c_{s}(t,z) - c_{P}^{*}(t,z)\right)$$
(11)

where  $\mu$  is sphericity, l a characteristic dimension of particles and can be defined as l = r/3, r is the mean particle radius,  $\rho_s$  is the solid density,  $D_i(T(t,z))$  corresponds to the overall diffusion coefficient and  $c_P^*(t,z)$  is a 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. [17], a linear equilibrium relationship (equation 12) can be used to find an equilibrium concentration of the solute in the fluid phase  $c_f^*(t, z)$  is based on concentration of the solute in the solid phase  $c_s(t, z)$ 

$$c(t,z) = \frac{k_n(T(t,z), P(t))q^*(t,z)}{(12)}$$

The volumetric partition coefficient  $k_p(T(t,z), P(t))$  behaves 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 [18], the term  $k_p(T(t,z), P(t))$  can be expressed as the function of mass partition factor  $k_m(T(t,z))$ .

$$k_{m}(T(t,z)) = \frac{k_{p}(T(t,z), P(t))\rho_{s}}{\rho(T(t,z), P(t))}$$
(13)

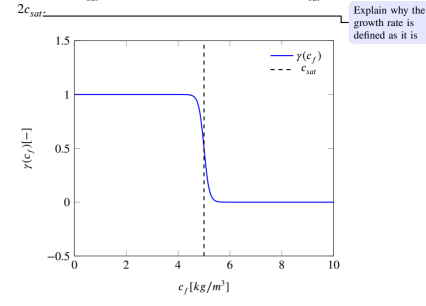
Equation 14 represents of the kinetic term according to Reverchon [1]

$$r_e(t,z) = -\frac{\frac{D_i(T(t,z), P(t))}{\mu l^2} \left(c_s(t,z) - \frac{\rho_s}{\frac{k_m(T(t,z))\rho(T(t,z), P(t))}{\rho(T(t,z), P(t))}}c_f(t,z)\right)$$
(14)

The above model does not take into account the saturation of fluid, which can be introduced by multiplying the gradient by a function  $\gamma(c_f)$  (equation 15).  $\gamma(c_f)$  describe the reverse logistic function, which is equal to unity below the  $c_{sat}$ , the saturation concentration, and equal to zero, above the  $c_{sat}$ . The  $\gamma(c_f)$  for  $c_{sat}=5$  is shown on figure 7.

$$\gamma(c_f) = \frac{1}{1 + \exp(-k_{sat} (c_f - c_{sat}))}$$
 (15)

where  $k_{sat}$  is the growth rate, and it is defined as  $k_{sat}$  =



**Figure 7:**  $\gamma(c_f)$  function under assumption of  $c_{sat} = 5[kg/m^3]$ 

The final form of the extraction kinetic equation is given by equation 16.

$$r_e(t,z) = -\frac{D_i(T(t,z), P(t))}{\mu l^2} \gamma(c_f) \left( c_s(t,z) - \frac{\rho_s}{k_m(T(t,z))\rho(T(t,z), P(t))} c_f(t,z) \right)$$
(16)

#### 2.4.5. Heat balance

The heat equation was introduced in the previous chapter through equation 5 or

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

For real gases the internal energy is defined as

$$de = C_v dT - \left| P - T \left( \frac{\partial P}{\partial T} \right)_{V_m} \right| dv_m \tag{17}$$

If a gas is consider to be perfectly caloric ( $e=C_vT$ ), then the energy equation can be written in form of temperature. The perfectly caloric gas can be seen as the special case of a real gas, where the second term of the equation 17 goes to zero.

For real gases it is complicated to write the heat balance in terms of temperature, but it be used directly in the form of internal energy, as it is given by equation 5. In such a case, the temperature (which is used as the input to some other functions) needs to be recover from the internal energy. A relation for the internal energy can be obtained from an equation of state. For Peng-Robinson, such a relation is given by equation 18.

$$e = C_v T + \frac{a\left(\alpha - T\frac{d\alpha}{dT}\right)}{2\sqrt{2}b} \ln\left[\frac{1 + b\left(1 - \sqrt{2}\rho\right)}{1 + b\left(1 + \sqrt{2}\rho\right)}\right]$$
(18)

To solve equation 18, values of temperature, pressure and density needs to be know. If an equation of state is introduce, then only 2 out 3 variables needs to be know as the third one can be calculated, this can be represented as follow

$$e(t,z)=e(T(t,z),P(t),\rho(t,z))=e(T(t,z),P(t),\rho(T(t,z),P(t)))$$

If the value of internal energy e is know from time evolution of the energy equation (5), and pressure is know from measurement, then the temperature can be reconstructed. A roootfinder can be used to find a value of temperature, which minimize the difference between value of internal energy coming from the time evolution and the output from equation 18. Such a procedure allow to find local temperature along spatial direction z and needs to be repeated every time-step.

Another way to express the energy equation is to introduce enthalpy  $e = h - P_0/\rho$ . By introducing the definition of enthalpy, the energy equation becomes

$$\frac{\partial \left(\rho_f h A_f\right)}{\partial t} - \frac{\partial \left(P_0 A_f\right)}{\partial t} + \frac{\partial \left(\rho_f h A_f v\right)}{\partial z} - \frac{\partial}{\partial z} \left(k \frac{\partial T}{\partial z}\right) \tag{19}$$

The main advantage of this formulation is the presence of term  $\partial P_0/\partial t$ , which allow to directly affect the system through the change of thermodynamic pressure (which is a control variable). If an equation of state is known, the temperature can to be recovered from the enthalpy. The enthalpy is related to the pressure and temperature through the following equation:

$$h(t,z) = h(T(t,z), P(t), \rho(t,z)) = h(T(t,z), P(t), \rho(T(t,z), P(t)))$$
 (20)

If the value of enthalpy is know from the time evolution and pressure can be measured, then the equation 20 can be solved for temperature to recover the temperature profile. By applying the departure functions to Peng-Robinson equation of state, the relation 20 can be expressed directly through equation 21 as presented in A.1.4 or given by Gmehling et al.

$$h_{T,P} - h_{T,P}^{\text{ideal}} = RT_C \left[ T_r(Z-1) - 2.078(1+\kappa) \sqrt{\alpha} \ln \left( \frac{Z+2.414B}{Z-0.414B} \right) \right] \tag{21}$$

The equation 21 requires an reference sate, which in this case is assumed to be  $T_{ref} = 298.15[K]$  and  $P_{ref} = 1.01325[bar]$ .

Add short section about boundary conditions and comment on the used numerical methods

### 2.4.6. Extraction yield

The efficiency of the process (the yield) is calculated according to equations 22 to 23, which evaluate the mass of solute at the exit of the extraction unit and sums it. The integral form of the measurement equation can be transformed into the differential form, and augmented with model equations to be solved simultaneously.

$$y(t)[kg] = \int_{t_0}^{t_f} \frac{F(t)}{\rho(t,z)} \left[ \frac{kg}{s} \left( \frac{kg}{m^3} \right)^{-1} \right] c_f(t,z) \Big|_{z=L} \left[ \frac{kg}{m^3} \right] dt[s]$$

$$\frac{dy}{dt} \left[ \frac{kg}{s} \right] = \frac{F(t)}{\rho(t,z)} \left[ \frac{kg}{s} \left( \frac{kg}{m^3} \right)^{-1} \right] c_f(t,z) \Big|_{z=L} \left[ \frac{kg}{m^3} \right]$$
 (23)

2.5. Parameter estimation

Conceptually, the unobservable error  $\epsilon(t)$  is added to the deterministic model output, y(t) (equation 22), to give the observable dependent variable Y(t) (for example results of an experiments). For discrete observations:

The same symbol as void fraction

$$Y(t_i) = y(t_i) + \epsilon(t_i)$$

and for continuos variables:

$$Y(t) = y(t) + \epsilon(t)$$

The objective in parameter estimation is to obtain the "best" estimate of  $\theta$  based on the continuous observations Y(t) or the discrete observations  $Y(t_i)$ . The above equation can be written in general form as:

$$Y(t_i) = y(\theta, t_i) + \epsilon(t_i)$$
(24)

where  $\theta$  is set of parameters to be estimated from the data.  $\theta$  is a subset of parameter space  $\Theta$ , which contain all parameters of the model.

Because of the difficulty of obtaining analytical solutions to the deterministic process model, experiments have been arranged whereby the vector of derivatives  $dY(t_i)/dt$  is measured rather than  $Y(t_i)$  itself. In such cases, it is assumed that the unobservable error is added to the deterministic derivative  $dy(\theta, t_i)/dt$  (equation 23) as follows:

$$\frac{dY(t_i)}{dt} = \frac{dy(\theta, t_i)}{dt} + \epsilon(t_i)$$

If unobservable error in the first observation is designated as  $\epsilon_1$ , the second error observation  $\epsilon_2'$  includes  $\epsilon_1$  plus a random component introduced side from  $\epsilon_1$ , or  $\epsilon_2' = \epsilon_1 + \epsilon_2$ . The error in the third observation is  $\epsilon_3' = \epsilon_1 + \epsilon_2 + \epsilon_3$ , and so forth. Mandel [20] distinguished between the usually assumed type of independent measuring error in the dependent variable and a 'cumulative' or interval error in which each new observation includes the error of the previous observations. Cumulative errors, arising because of the fluctuations as a function of time in the process itself due to small changes in operating, are not independent - only the differences in measurement from one period to the next are independent.

#### 2.5.1. Least squares estimation

A least squares parameter estimation does not require prior knowledge of the distribution of unobservable errors, yield unbiased estimates, and results in the minimum variance among all linear unbiased estimators. If the observations Y for the model responses are continuous functions of time from t=0 to  $t=t_f$ , the Markov (or "rigorous least squares") criterion is to minimize:

$$\phi = \frac{1}{2} \int_0^{t_f} [Y(t) - y(\theta, t)]^T \Gamma^{-1} [Y(t) - y(\theta, t)] dt$$
 (25)

$$\hat{\theta} = \arg\min_{\theta \in \Theta} \phi \tag{26}$$

where  $\Gamma$  is the covariance matrix (or perhaps a matrix of appropriate weights),  $\hat{\theta}$  is the parameter estimate,  $\Theta$  is the parameter space, and  $\phi$  is the time integrated value of the error squared ("integral squared error"). If the observations

"For high pressures, the influence of the intermolecular forces on the enthalpy has to be taken into account, usually by applying a cubic equation of state. In most cases, these forces are attractive, so that additional energy is necessary to move the molecules away from each other. that is, to lower the density. If this energy is not added, the substance cools down when it is expanded" (Gmehling et al. [19]). So, pressure reduction cause gas expansion and temperature drop. Give reference

for measurement

function

are made at discrete instants of time,  $t_i$ , i = 1, 2, ..., n, the Markov criterion is to minimize:

$$\phi = \frac{1}{2} \sum_{i=1}^{n} \left[ Y(t_i) - y(\theta, t_i) \right]^T \Gamma^{-1} \left[ Y(t_i) - y(\theta, t_i) \right]$$
 (27)

$$\hat{\theta} = \arg\min_{\theta = 0} \phi \tag{28}$$

If  $\Gamma$  is a diagonal matrix,  $\phi$  becomes a "weighted least squares" criterion; if  $\Gamma = \sigma^2 \mathbf{I}$  where  $\sigma$  is a standard deviation,  $\phi$  is the "ordinary least squares" criterion.

### 2.5.2. Maximum likelihood estimation

Maximum likelihood estimation (MLE) is a method of estimating the parameters of an assumed probability distribution, given some observed data. This is achieved by maximizing a likelihood function so that, under the assumed statistical model, the observed data is most probable. The MLE has the desirable characteristics of asymptotic efficiency and normality. Each time it has been associated with the (joint) normal distribution because of mathematical convenience. Consider the joint probability density function (the likelihood function)  $p\left(\theta|y(t_1),y(t_2),...,y(t_n)\right)$  for parameter  $\theta$ . If a maximum of this function over all choices of  $\theta$  can be found, the estimates so obtained are maximum likelihood estimates. The conditions at the maximum can be evolved incorporating prior information as follows.

The posterior probability density  $p(\theta|y)$  can be expressed as the ratio of two probability densities if we make use of the analogue for continuous variables of Equation 109:

$$p\left(\theta | y(t_n), ..., y(t_1)\right) = \frac{p\left(\theta, y(t_n), ..., y(t_1)\right)}{p\left(y(t_n), ..., y(t_1)\right)} \tag{29}$$

The numerator of the right-hand side od Equation 29 using Equation 110a becomes

Notation is a bit unclear

$$p(\theta, y(t_n), ..., y(t_1)) = p(y(t_n)|\theta, y(t_{n-1}), ..., y(t_1))$$

$$\cdot p(\theta, y(t_{n-1}), ..., y(t_1))$$
(30)

These operations can be continued repetitively until we get

$$p(\theta, y(t_n), ..., y(t_1)) = p(\theta) \prod_{i=1}^{n} p(y(t_i)|\theta, y(t_{i-1}), ..., y(t_1))$$
(31)

Examination of Equation 24 shows that  $Y(t_i)$  depends only on  $t_i$ ,  $\theta$  and  $\varepsilon(t_i)$  and is not conditioned by any previous measurement. Consequently, we can write

$$p(y(t_i)|\theta, y(t_{i-1}), ..., y(t_1)) = p(y(t_i)|\theta)$$
(32)

provided Equation 24 is observed as a constraint. The desired joint conditional probability function is thus

$$p(\theta|y(t_n), ..., y(t_1)) = \frac{p(\theta) \prod_{i=1}^{n} p(y(t_i)|\theta)}{p(y(t_n), ..., y(t_1))}$$
(33)

We can get rid of the evidence term  $p\left(y(t_n),...,y(t_1)\right)$  because it's constant with respect to the maximization. Moreover, if we are lacking a prior distribution over the quantity we want to estimate, then  $p(\theta)$  can be omitted. In such a case:

$$p\left(\theta|y(t_n),...,y(t_1)\right) = \prod_{i=1}^n p\left(y(t_i)|\theta\right) = \prod_{i=1}^n L\left(\theta|y(t_i)\right)$$
(34)

By collecting a values of y and selecting the values of  $\theta$  that maximize the likelihood function  $L(\theta|y)$ , a function described in chapter A.3 in connection with Bayes' theorem. Such estimators,  $\hat{\theta}$ , are known as maximum likelihood estimators. In effect, the methods selects those values of  $\theta$  that are at least as likely to generate the observed sample as any other set of values of the parameters if the probability density of y were to be extensively simulated through use of the probability density  $p(y|\theta)$ . In making a maximum likelihood estimate, we assume the form of the probability density and that all possible values of  $\theta$  are equally likely before experimentation.

□ Needed?

needed?

Is the reference it previous chapter

The likelihood function for the parameters based on several observations is the product of the individual functions if the observations are independent.

$$L(\theta|y(t_n), ..., y(t_1)) = \prod_{i=1}^{n} L(\theta|y(t_i))$$
  
=  $p(y(t_1)|\theta) p(y(t_2)|\theta) ... p(y(t_n)|\theta)$  (35)

In choosing as estimates of  $\theta$  the values that maximize L for the given values  $(y(t_i))$ , it turns out that it is more convenient to work with the  $\ln L$  than with L itself:

$$\ln L = \ln p\left(y(t_1)|\theta\right) + \ln p\left(y(t_2)|\theta\right) + \dots + \ln p\left(y(t_n)|\theta\right) = \sum_{i=1}^n \ln p\left(y(t_i);\theta\right)$$
(36)

By assuming that the conditional distribution of  $\bar{Y}_i$ , given  $y_i$ , is normal, then we form the likelihood function based on the probability density:

$$p\left(\theta, \sigma | y(t_n), ..., y(t_1)\right) = \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi}\sigma} \exp\left[\frac{1}{2\sigma} \left(Y(t_i) - y(\theta, t_i)\right)^2\right]$$

$$L\left(\theta, \sigma | y(t_n), ..., y(t_1)\right) = \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi}\sigma} \exp\left[\frac{1}{2\sigma} \left(Y(t_i) - y(\theta, t_i)\right)^2\right]$$
(37)

where  $\sigma$  is the variance

By taking the natural logarithm of the Equation 37, the final form of the objective function can be obtained:

$$\ln L = -\frac{n}{2} \left( \ln \sqrt{2\pi} + \ln \sigma^2 \right) - \frac{\sum_{i=1}^{n} \left[ Y(t_i) - y(\theta, t_i) \right]^2}{2\sigma^2}$$
 (38)

The parameter estimation problem can be formulated as follow:

$$\hat{\theta}_{MLE} = \arg\max_{\sigma,\theta \in \Theta} \ln L = \arg\max_{\sigma,\theta \in \Theta} p(\theta|y)$$
 subject to 
$$\dot{x} = f(t,x,\theta)$$
 
$$\dot{\theta} = 0$$
 (39)

x has not been

Based on the first order optimality condition, the  $\ln L$  can be maximized with respect to the vector  $\theta$  by equating to zero the partial derivatives of  $\ln L$  with respect to each of the parameters:

$$\frac{\partial \ln L}{\partial \theta} = \frac{\partial \sum_{i=1}^{n} \ln p \left( y(t_i) | \theta \right)}{\partial \theta} = 0 \tag{40}$$

Solution of Equations 40 yield the desired estimates  $\hat{\theta}$ . For some models, these equations can be explicitly solved for  $\hat{\theta}$  but in general no closed-form solution to the maximization problem is known or available, and an MLE can only be found via numerical optimization.

## 2.6. Experimental work

The close the optimization problem given by set of equations 39, it is require to know the dataset Y(t). In this case, the dataset have been obtained by performing extraction of caraway oil from caraway seeds. The caraway seeds have been obtained from the company Caraway Finland during 2022 growing season. The caraway seeds have been pre-treated with Retsch SM 300 cutting mill to decrease the particle size to 1mm and break the outer shell of the seeds. Later, the moisture content was analysed to ensure that it does not have a negative effect on the extraction process. An Infrared Moisture Analyser was used to investigate the sample. The moisture content after grinding was 4.83%. The grind material density was measured by pyncometery and its result can be found in appendix A.4

Should Biorukki be mentioned?

Should be mentioned?

Double check dimension The unit used for the experimental work is 10 litre extractor with inner diameter around 15 cm and 60 cm height. Four experiments have been performed in that unit at following operating conditions:  $40^{\circ}C/200$  bar,  $50^{\circ}C/200$  bar,  $40^{\circ}C/300$  bar and  $50^{\circ}C/300$  bar. 1 kg of grind material was weighted and placed in the extraction chamber. Due to some technical limitations, the amount of solid material fro extraction was selected to be 1 kg per batch, which is not sufficient to fill the whole chamber.

After the material was loaded into chamber, the extractor is pre-heated to desired temperature. The next step is to close the outlet line coming out of the extractor and fill the extraction chamber with  $CO_2$ . The  $CO_2$  is pumped and compressed until the operating pressure is achieved. When the operating temperature and pressure are obtained, the outlet in is open to allow the solvent flow through the system. The solvent extract essential oils from the fixed bed of solid material and the mixture  $CO_2$ -oil flow from the extractor to separator. The separator was operating at  $50^{\circ}C$  and 50 bar. At this condition the  $CO_2$  is gasified and leave from the top, while oil stays in the liquid form and it is collected at the bottom of separator. Every 5 minutes, the oil is drained from the separator and its weight is measured. The amount of oil collect during a batch is an output of the experiment and it is used as data for parameter estimation. The extraction time for each batch was 150 minutes, counted from the opening of the extractor outlet port.

The result of the experiment are presented in the table 1.

## 3. Results

Some results of the parameter estimations

### 4. Conclusions

	40°C/200 bar	50°C/200 bar	40°C/300 bar	50°C/300 bar
Time min	Mass g	Mass g	Mass g	Mass g
0	0.0	0.0	0.0	0.0
5	1.7	1.8	1.1	1.2
10	5.7	4.7	4.7	4.5
15	10.9	9.5	10.8	11.3
20	17.8	9.5 15.1	20.6	20.6
25	24.9	21.3	32.1	30.6
30		26.4	32.1 40.5	
30 35	30.3 34.0	20.4 29.8	40.5 46.7	37.8 43.3
40	37.5	33.1	50.5	47.5
45	40.1	35.6	53.1	50.8
50	43.3	38.7	55.5	53.7
55	45.5	41.5	57.6	56.5
60	47.6	43.5	59.3	58.1
65	49.2	45.9	60.9	59.6
70	50.8	48.3	62.2	61.1
75	52.6	50.8	63.7	62.7
80	54.2	52.9	64.9	64.4
85	55.1	55.3	66.6	65.8
90	56.2	57.2	67.6	66.7
95	57.3	58.3	68.9	67.4
100	58.3	59.4	69.7	68.2
105	59.4	60.4	70.6	69.1
110	60.2	61.6	71.2	70.1
115	61.4	62.7	71.9	70.9
120	62.5	63.9	72.3	71.6
125	63.3	64.7	72.8	72.3
130	64.0	65.4	73.2	72.9
135	64.8	65.9	73.6	73.5
140	65.5	66.4	74	74
145	66.2	67.1	74.3	74.5
150	66.8	67.7	74.6	74.9
_50	1 23.0			

**Table 1** Yield data

## Parameter estimation

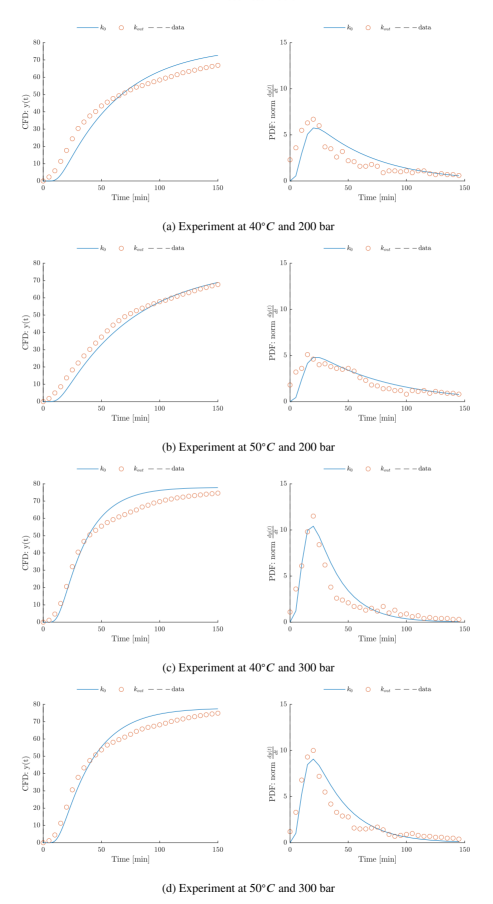


Figure 8: Results of parameter fitting, without estimation of the initial state

## Parameter estimation

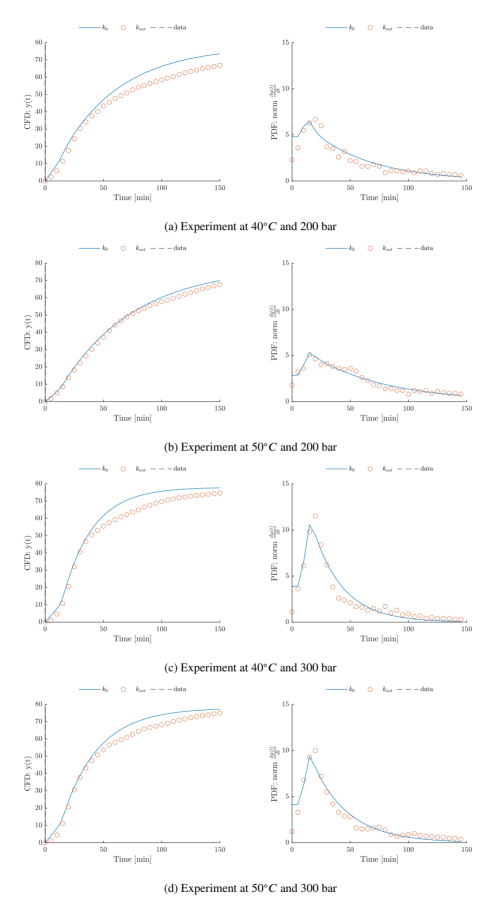


Figure 9: Results of parameter fitting, with estimation of the initial state

#### References

- E. Reverchon. Mathematical modeling of supercritical extraction of sage oil. <u>AIChE Journal</u>, 42(6):1765–1771, jun 1996. doi: 10.1002/aic.690420627.
- [2] Zhen Huang, Xiao han Shi, and Wei juan Jiang. Theoretical models for supercritical fluid extraction. <u>Journal of Chromatography A</u>, 1250: 2–26, aug 2012. doi: 10.1016/j.chroma.2012.04.032.
- [3] Ernesto Reverchon, Giorgio Donsi, and Libero Sesti Osseo. Modeling of supercritical fluid extraction from herbaceous matrices. <u>Industrial</u> & Engineering Chemistry Research, 32(11):2721–2726, nov 1993. doi: 10.1021/ie00023a039.
- [4] H. Sovova. Rate of the vegetable oil extraction with supercritical co2. modelling of extraction curves. <u>Chemical Engineering Science</u>, 49 (3):409–414, 1994. doi: 10.1016/0009-2509(94)87012-8.
- [5] Ding-Yu Peng and Donald B. Robinson. A new two-constant equation of state. <u>Industrial & Engineering Chemistry Fundamentals</u>, 15(1): 59–64, feb 1976. doi: 10.1021/i160057a011.
- [6] J Elliott. Introductory chemical engineering thermodynamics. Prentice Hall, Upper Saddle River, NJ, 2011. ISBN 9780136068549.
- [7] R. M. Pratt. Thermodynamic properties involving derivatives: Using the peng-robinson equation of state, chemical engineering education. <u>CHEMICAL ENGINEERING EDUCATION</u>, 35:112–139, 2001. ISSN 0009-2479. URL https://www.tib.eu/de/suchen/id/BLSE% 3ARN995457161
- [8] G. G. Simeoni, T. Bryk, F. A. Gorelli, M. Krisch, G. Ruocco, M. Santoro, and T. Scopigno. The widom line as the crossover between liquid-like and gas-like behaviour in supercritical fluids. <u>Nature Physics</u>, 6(7):503–507, jun 2010. doi: 10.1038/nphys1683.
- [9] Daniel Banuti. The latent heat of supercritical fluids. <u>Periodica Polytechnica Chemical Engineering</u>, 63(2):270–275, jan 2019. doi: 10.3311/ppch.12871.
- [10] W. Sheng, G. J. Chen, and H. C. Lu. Prediction of transport properties of dense gases and liquids by the peng-robinson (PR) equation of state. <u>International Journal of Thermophysics</u>, 10(1):133–144, jan 1989. doi: 10.1007/bf00500714.
- [11] A. Fenghour, William A. Wakeham, and V. Vesovic. The viscosity of carbon dioxide. <u>Journal of Physical and Chemical Reference Data</u>, 27 (1):31–44, jan 1998. doi: 10.1063/1.556013.
- [12] Arno Laesecke and Chris D. Muzny. Reference correlation for the viscosity of carbon dioxide. <u>Journal of Physical and Chemical</u> Reference Data, 46(1):013107, mar 2017. doi: 10.1063/1.4977429.
- [13] M. L. Huber, E. A. Sykioti, M. J. Assael, and R. A. Perkins. Reference correlation of the thermal conductivity of carbon dioxide from the triple point to 1100 k and up to 200 MPa. <u>Journal of Physical and Chemical Reference Data</u>, 45(1):013102, mar 2016. doi: 10.1063/1. 4940892.
- [14] John D. Anderson. <u>Computational fluid dynamics the basic with applications</u>. McGraw-Hill, 1995. ISBN 9780071132107.
- [15] Stefan Schreier. Compressible flow. Wiley, 1982. ISBN 047105691X.
- [16] Pierre-Louis Lions. Mathematical Topics in Fluid Mechanics: Volume 1 Incompressible Models. Oxford University Press, 2013. ISBN 9780199679218.
- [17] N. R. Bulley, M. Fattori, A. Meisen, and L. Moyls. Supercritical fluid extraction of vegetable oil seeds. <u>Journal of the American</u> Oil Chemists' Society, 61(8):1362–1365, aug 1984. doi: 10.1007/ bf02542243.
- [18] M. Spiro and M. Kandiah. Extraction of ginger rhizome: partition constants and other equilibrium properties in organic solvents and in supercritical carbon dioxide. <u>International Journal of Food Science</u> & Technology, 25(5):566–575, jun 2007. doi: 10.1111/j.1365-2621. 1990.tb01116.x.
- [19] Jürgen Gmehling, Michael Kleiber, Bärbel Kolbe, and Jürgen Rarey. Chemical Thermodynamics for Process Simulation. Wiley, mar 2019. doi: 10.1002/9783527809479.
- [20] John Mandel. Fitting a straight line to certain types of cumulative data. <u>Journal of the American Statistical Association</u>, 52(280):552– 566, dec 1957. doi: 10.1080/01621459.1957.10501413.

[21] David Mautner Himmelblau. Process analysis by statistical methods, 1970. [by] David M. Himmelblau., Includes bibliographical references

## A. Appendix

## A.1. Thermodynamic

# A.1.1. Equation of state and properties of the fluid phase

We consider equations of state in the general form  $P(t)V_{\text{CO}_2}[T(t,z), P(t)] = ZRT(t,z)$ , where  $V_{\text{CO}_2}$  denotes the molar volume of  $\text{CO}_2$ , Z represents its compressibility factor, and R is the universal gas constant. More specifically, we are interested in these equations because of the possibility to express the compressibility Z as an explicit function of temperature and pressure. This is the case when Z is obtained as one of the physically meaningful roots of a polynomial equation, like the Peng-Robinson's and stuff with REFS to equations of state.

time, due to its dependence on temperature and pressure, and it is expressed in terms of the set of physical constants  $\varphi_{\rho_{\rm E}} = \left(R, M_{\rm CO_2}, T_{\rm CO_2}^c, P_{\rm CO_2}^c, \omega_{\rm CO_2}\right)$ .

## A.1.3. Heat capacity of the fluid phase

The specific heat  $C_p^{\rm F}$  can be calculated from the equation of state, again under the assumption that the fluid phase consists of pure carbon dioxide and that the specific heat of real fluids can be calculated from an ideal contribution plus a residual term REF. In the following, we report only the main steps in the derivation of  $C_p^{\rm CO_2}[T(t,z),P(t)]$ : Step-by-step derivations using the Peng-Robinson's equation are given.

Add step-by-step derivation

Add REF

At given temperature and pressure, for CO<sub>2</sub> we have

reference to Cardano formula In Peng-Robinson's equation, the compressibility  $\overline{Z}[T(t,z), P(t)] C_v^{CO_2}[T(t,z), P(t)] = C_v^{I}[T(t,z), P(t)] + C_v^{R}[T(t,z), P(t)]$ ; solves the third-order polynomial equation

$$Z^{3} - [1 - B[T(t, z), P(t)]]Z^{2}$$

$$+ [A[T(t, z), P(t)] - 2B[T(t, z), P(t)] - 3B[T(t, z), P(t)]^{2}]Z = 0$$
(41)

where A[T(t,z), P(t)] and B[T(t,z), P(t)] are functions of time and space defined on the attraction parameter,  $a[T(t,z)] = a_{\text{CO}_2}^c \alpha[T(t,z)]$  with  $a_{\text{CO}_2}^c \approx 0.45724 R^2 T_{\text{CO}_2}^c ^2 / P_{\text{CO}_2}^c$ , and the repulsion parameter,  $b_{\text{CO}_2} \approx 0.07780 R T_{\text{CO}_2}^c / P_{\text{CO}_2}^c$ , both functions of the critical temperature  $T_{\text{CO}_2}^c$  and pressure  $T_{\text{CO}_2}^c$ . Specifically, we have

$$A[T(t,z), P(t)] = \frac{\alpha [T(t,z)] a_{\text{CO}_2}^c P(t)}{R^2 T^2(t,z)};$$
(42a)

$$B[T(t,z), P(t)] = \frac{b_{\text{CO}_2}P(t)}{RT(t,z)}.$$
(42b)

The quantity  $\alpha [T(t,z)] = \left[1 + \kappa_{\text{CO}_2} \left[1 - \sqrt{T(t,z)/T_{\text{CO}_2}^c}\right]\right]^2$ , with constant  $\kappa_{\text{CO}_2} = 0.37464 + 1.54226\omega_{\text{CO}_2} - 0.26992\omega_{\text{CO}_2}^2$ , is a dimensionless correction term defined on the acentric factor  $\omega_{\text{CO}_2} = 0.239$  of CO<sub>2</sub> molecules.

By denoting the physical constants as  $\varphi_Z = \left(R, T_{\text{CO}_2}^c, P_{\text{CO}_2}^c, \kappa_{\text{CO}_2}^{\text{at}}\right)$ , on the pressure P(t), we have we obtain a spatio-temporal representation of the compressibility  $\overline{Z}\left[T(t,z), P(t) \mid \varphi_Z\right]$  with its complete set of functional dependencies and parameters.

## A.1.2. Density of the fluid phase

The density  $\rho_{\rm F}$  of the fluid phase is assumed to be equal to the density of solvent, at given temperature and pressure. Because temperature T(t,z) of the fluid phase is a modelled variable, we allow for the density to vary along the bed and in time. From an equation of state of the form  $P(t)V_{\rm CO}$ , [T(t,z),P(t)] = ZRT(t,z), we get

$$\rho_{F}\left[T(t,z), \mathbf{P}(t) \mid \varphi_{\rho_{F}}\right] = \frac{\mathbf{P}(t)\mathbf{M}_{CO_{2}}}{\mathbf{R}T(t,z)\overline{\mathbf{Z}}\left[T(t,z), \mathbf{P}(t) \mid \varphi_{Z}\right]},\tag{43}$$

where  $M_{\text{CO}_2}$  denotes the molar mass of  $\text{CO}_2$  and  $\overline{Z}[T(t,z), P(t)]$  is the compressibility factor that solves Eq. (41). The density of the fluid is thus a function of space and

(1) = (0) = (1) = (

$$C_{p}^{\text{CO}_{2}}[T(t,z), P(t)] = \underbrace{C_{p}^{1}[T(t,z), P(t)]}_{\text{Eq. (45)}} + \underbrace{C_{p}^{R}[T(t,z), P(t)]}_{\text{Eq. (46)}}.$$
(44a)

 $C_v^{\text{CO}_2}[T(t,z), P(t)]$  and  $C_p^{\text{CO}_2}[T(t,z), P(t)]$  are the specific heat of  $\text{CO}_2$  at constant volume and pressure, respectively.  $C_v^{\text{I}}[T(t,z), P(t)]$  and  $C_p^{\text{I}}[T(t,z), P(t)]$ , with  $C_p^{\text{I}}(T(t,z)) - C_v^{\text{I}}(T(t,z)) = R$ , are the specific heat of an ideal gas at constant volume and pressure.  $C_v^{\text{R}}[T(t,z), P(t)]$  and  $C_p^{\text{R}}(T(t,z), P(t))$  are the correction terms.

For  $CO_2$  (), we have the ideal gas contribution to the specific heat at constant P(t), as function of T(t,z),

$$C_{p}^{I}[T(t,z),P(t)] = C_{p_0} + C_{p_1}T(t,z) + C_{p_2}T^2(t,z) + C_{p_3}T^3(t,z)$$
 (45)

where the coefficients of the expansion are  $C_{P0} = 4.728$ ,  $C_{P1} = 1.75 \times 10^{-3}$ ,  $C_{P2} = -1.34 \times 10^{-5}$ , and  $C_{P3} = 4.10 \times 10^{-9}$ . For the correction term  $C_p^R$  [T(t, z)P(t)] at constant pressure P(t), we have

$$C_{p}^{R}[T(t,z), P(t)] = \underbrace{C_{v}^{R}[T(t,z), P(t)]}_{\text{Eq. (50)}} + T(t,z) \underbrace{\left(\frac{\partial P(t)}{\partial T}\right)_{V_{\text{CO}_{2}}(t,z)}}_{\text{Eq. (49)}} \underbrace{\left(\frac{\partial V_{\text{CO}_{2}}[T(t,z), P(t)]}{\partial T}\right)_{P(t)}}_{\text{Eq. (47)}} - R. \quad (46)$$

The braced terms are obtained from the chosen equation of state P(t)V[T(t, z), P(t)] = Z[T(t, z), P(t)]RT(t, z).

For the partial derivative of the volume with respect to temperature T at constant pressure P(t), we have

$$\left(\frac{\partial V_{\text{CO}_2}\left[T(t,z),P(t)\right]}{\partial T}\right)_{P(t)} = \frac{\frac{Z\left[T(t,z),P(t)\right]R}{P(t)} + \frac{RT(t,z)}{P(t)} \underbrace{\left(\frac{\partial Z\left[T(t,z),P(t)\right]}{\partial T}\right)_{P(t)}}_{\text{Eq. (48)}}$$

$$(47)$$

with partial derivative of the compressibility factor with respect to temperature T at constant pressure P(t)

$$\left(\frac{\partial Z\left[T(t,z),P(t)\right]}{\partial T}\right)_{P(t)} = \left(\frac{\partial \frac{P(t)V_{\text{CO}_2}\left[T(t,z),P(t)\right]}{RT(t,z)}}{\partial T}\right)_{P(t)} \tag{48}$$

Similarly, for the partial derivative of the pressure with respect to temperature at constant volume, we have

$$\left(\frac{\partial P(t)}{\partial T}\right)_{V_{\text{CO}_2}(t,z)} = \left(\frac{\partial \frac{ZRT(t,z)}{V_{\text{CO}_2}\left[T(t,z),P(t)\right]}}{\partial T}\right)_{V(t,z)} \tag{49}$$

The residual specific heat at constant volume is obtained, by definition, by using the residual internal energy

$$C_{v}^{R}\left[T(t,z), \mathbf{P}(t)\right] = \left(\frac{\partial U^{R}\left[T(t,z), \mathbf{P}(t)\right]}{\partial T}\right)_{V(t,z)}.$$
(50)

By denoting the physical constants as  $\varphi_{C_p^F} = \left( R, M_{CO_2}, T_{CO_2}^c, P_{CO_2}^c, \frac{\partial}{\partial t} \iint_{\mathcal{V}_f} \varphi \right) d\mathcal{V}_f + \iint_S \rho_f \mathbf{V} \cdot d\mathbf{S} = 0$ we get the spatio-temporal representation  $C_p^{\mathrm{F}}\left[T(t,z),P(t)\mid\varphi_{C_p^{\mathrm{F}}}\right]$ of the specific heat of the fluid phase.

## Change to my text $\neg$ A.1.4. Departure functions for enthalpy calculations

In thermodynamics, a departure function is defined for any thermodynamic property as the difference between the property as computed for an ideal gas and the property of the species as it exists in the real world, for a specified temperature T and pressure P. Common departure functions include those for enthalpy, entropy, and internal energy.

Departure functions are used to calculate real fluid extensive properties (i.e. properties which are computed as a difference between two states). A departure function gives the difference between the real state, at a finite volume or non-zero pressure and temperature, and the ideal state, usually at zero pressure or infinite volume and temperature.

For example, to evaluate enthalpy change between two points  $h(V_1, T_1)$  and  $h(V_2, T_2)$  we first compute the enthalpy departure function between volume  $V_1$  and infinite volume at  $T = T_1$ , then add to that the ideal gas enthalpy change due to the temperature change from  $T_1$  to  $T_2$ , then subtract the departure function value between  $V_2$  and infinite volume.

Departure functions are computed by integrating a function which depends on an equation of state and its derivative. The general form the enthalpy equation is given

$$\frac{H^{ig} - H}{RT} = \int_{V}^{\infty} \left[ T \left( \frac{\partial Z}{\partial T} \right)_{V} \right] \frac{dV}{V} + 1 - Z \tag{51}$$

The Peng-Robinson equation of state relates the three interdependent state properties pressure P, temperature T, and molar volume  $V_m$ . From the state properties  $(P, V_m, T)$ , one may compute the departure function for enthalpy per mole (denoted h) as presented by Gmehling et al. [19] or Elliott [6]:

$$h_{T,P} - h_{T,P}^{\text{ideal}} = RT_C \left[ T_r(Z-1) - 2.078(1+\kappa) \sqrt{\alpha} \ln \left( \frac{Z+2.414B}{Z-0.414B} \right) \right] \tag{52}$$

## A.2. Governing equations

#### A.2.1. Mass continuity

Following the work of Anderson [14], the governing equations for compressible fluid with non-uniform crosssection can be obtained. Let's assume that any properties of the flow are uniform across any given cross-section of an extractor. The variation of the cross-section might be an result of partial filling of an extractor or its irregular shape. In reality, such a flow is two-dimensional, because with the area changing as a function of z, in actuality there will be flow-field variations in both directions. The assumption of quasi-one-dimensional flow dictates that the flow properties are function of z only. The equations described by quasione-dimensional assumption hold: (1) mass conservation, (2) Newton's second law, and (3) energy conservation. To ensure that these physical principles are satisfied the modified governing equation can be derived. Let's start with the integral form of the continuity equation:

$$\frac{P_{\text{CO}}^{c}}{\partial t} \iiint_{\mathcal{V}_{c}} \rho_{f} d\mathcal{V}_{f} + \iint_{S} \rho_{f} \mathbf{V} \cdot \mathbf{dS} = 0$$
(53)

We apply this equation to the shaded control volume shown in Fig. 10.

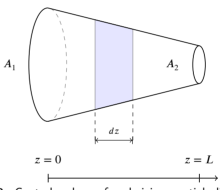


Figure 10: Control volume for deriving partial differential equation for unsteady, quasi-one-dimensional flow

This control volume is a slice of an extractor, where the infinitesimal thickens of the slice is dz. On the left side of the control volume, consistent with the quasi-onedimensional assumptions, the density, velocity, pressure and internal energy denoted by  $\rho_f$ , V, P, and e, respectively, are uniform over the are A. Similarly, on the right side of the control volume, the density, velocity, pressure, and internal energy  $\rho_f + d\rho_f$ , V + dV, P + dP, and e + de, respectively, are uniform over the area available for fluid phase  $A_f + dA_f$ . Applied to the control volume in Fig. 10, the volume integral in Eq. 53 becomes, in the limit as dz becomes very small,

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}_f} \rho_f d\mathcal{V}_f = \frac{\partial}{\partial t} \left( \rho_f A_f dz \right) \tag{54}$$

where A dz is the volume of the control volume in the limit of dz becoming vanishingly small. The surface integral

$$\iint_{S} \rho_{f} \mathbf{V} \cdot \mathbf{dS} = -\rho_{f} V A_{f} + (\rho_{f} + d\rho_{f})(V + dV)(A_{f} + dA_{f}) \tag{55}$$

where the minus sign on the leading term on the right-hand side is due to the vectors  $\mathbf{V}$  and  $\mathbf{dS}$  pointing in opposite directions over the left of the control volume, and hence the dot product is negative. Expending the triple product term

$$\iint_{S} \rho_{f} \mathbf{V} \cdot \mathbf{dS} = -\rho_{f} V A_{f} + \rho_{f} V A_{f} + \rho_{f} V d A_{f} + \rho_{f} A_{f} dV + \rho_{f} dV dA_{f} + V A_{f} d\rho_{f} + V d\rho_{f} dA + A_{f} d\rho_{f} dV + d\rho_{f} dV dA_{f}$$
(56)

In the limit as dz becomes very small, the terms involving products of the differential in Eq. 56, sych as  $\rho_f dV dA_f$ ,  $d\rho_f dV dA_f$ , go to zero much faster than those terms involving only one differential. Hence, all terms involving products of differentials can be dropped, yielding in the limit as dz becomes very small

$$\iint_{S} \rho_{f} \mathbf{V} \cdot \mathbf{dS} = \rho_{f} V dA_{f} + \rho_{f} A_{f} dV + V A_{f} d\rho_{f}$$
(57)

Substituting Eqs. 54 and 57 into 53, we have

$$\frac{\partial \left(\rho_f A_f\right)}{\partial t} + \frac{\partial \left(\rho_f A_f V\right)}{\partial z} = 0 \tag{58}$$

Above partial differential equation form of the continuity equation suitable for unsteady, quasi-one-dimensional flow. It ensures that mass is conserved for this mode of the flow. The  $A_f(z)$  is an arbitrary function, which describe change of the cross-section of an extractor. The function  $A_f(z)$  can be defined as  $A_f(z) = \mathbf{A}\varepsilon(z)$ , where  $\varepsilon$  is the bed porosity and  $\mathbf{A}$  is the cross-section of an empty extractor.

$$\frac{\partial \left(\rho_f \mathbf{A} \varepsilon(z)\right)}{\partial t} + \frac{\partial \left(\rho_f \mathbf{A} \varepsilon(z)V\right)}{\partial z} = 0 \tag{59}$$

The equation can be simplified by cancel out a constant A

$$\frac{\partial \left(\rho_f \varepsilon(z)\right)}{\partial t} + \frac{\partial \left(\rho_f \varepsilon(z)V\right)}{\partial z} = 0 \tag{60}$$

If so called superficial velocity is defined  $u = \varepsilon V$ , the mass continuity becomes

$$\frac{\partial \left(\rho_f \varepsilon(z)\right)}{\partial t} + \frac{\partial \left(\rho_f u\right)}{\partial z} = 0 \tag{61}$$

#### A.2.2. Transport of a species

The transport of a chemical species, in this case a solute, can be described by analogous equation to the Eq. 53 with additional terms on the right-hand side. The first term on the right-hand side describes that a substance goes from high density regions to low density regions and is based on the Fick's law  $\left(J_{diff} = D\frac{\partial C_f}{\partial z}\right)$ . The other term correspond for the mass transfer between solid and fluid phases, which is treated as a source term.

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}_f} C_f d\mathcal{V}_f + \iint_{S} C_f \mathbf{V} \cdot \mathbf{dS} = \iint_{S} J_{diff} \cdot \mathbf{n} \, \mathbf{dS} + \frac{\partial}{\partial t} \iiint_{\mathcal{V}_s} C_s d\mathcal{V}_s \tag{62}$$

Similarly to the continuity equation, in the limit as dz becomes very small

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}_f} C_f d\mathcal{V}_f = \frac{\partial}{\partial t} \left( C_f A_f dz \right) \tag{63}$$

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}} C_s d\mathcal{V}_s = \frac{\partial}{\partial t} (C_s A_s dz)$$
 (64)

The surface integrals in the limit of dz becomes

$$\iint_{S} C_{f} \mathbf{V} \cdot \mathbf{dS} = C_{f} V dA_{f} + C_{f} A_{f} dV + V A_{f} dC_{f}$$
 (65)

From the Divergence theorem in multi-variable calculus, we have

$$\iint_{S} J_{diff} \cdot \mathbf{n} \, d\mathbf{S} = \iiint_{\mathcal{V}_{f}} \nabla J_{diff} dV_{f} = \nabla \iiint_{\mathcal{V}_{f}} J_{diff} dV_{f} = \nabla \left( J_{diff} A_{f} dz \right)$$
(66)

By substituting the equations derived above into Eq. 62 we obtain

$$\frac{\partial \left(C_f A_f\right)}{\partial t} + \frac{\partial \left(C_f A_f V\right)}{\partial z} = \frac{\partial \left(C_s A_s\right)}{\partial t} + \frac{\partial \left(J_{diff} A_f\right)}{\partial z} \tag{67}$$

By defining  $A_f = A \cdot \varepsilon$ ,  $A_s = A \cdot (1 - \varepsilon)$  and  $u = V \cdot \varepsilon$ , and assuming that A is constant, the above equation becomes

$$\frac{\partial \left(C_{f}\varepsilon\right)}{\partial t} + \frac{\partial \left(C_{f}u\right)}{\partial z} = \frac{\partial \left(C_{s}(1-\varepsilon)\right)}{\partial t} + \frac{\partial \left(J_{diff}\varepsilon\right)}{\partial z} \tag{68}$$

By expanding above equation, splitting variable and assuming that  $\frac{\partial \varepsilon}{\partial t} = 0$  we get

$$\frac{\partial C_f}{\partial t} + \frac{u}{\varepsilon} \frac{\partial C_f}{\partial z} + \frac{C_f}{\varepsilon} \frac{\partial u}{\partial z} = \frac{1 - \varepsilon}{\varepsilon} \frac{\partial C_s}{\partial t} + \frac{D}{\varepsilon} \frac{\partial C_f}{\partial z} \frac{\partial \varepsilon}{\partial z} + \frac{\partial}{\partial z} \left( D \frac{\partial C_f}{\partial z} \right)$$
(69)

The equation can be further simplified if  $\frac{\partial u}{\partial z} = \frac{\partial \varepsilon}{\partial z} = D = 0$ , which corresponds to the assumptions of constant velocity along the bed( which might be a case of isothermal and low-mach number flow), constant porosity( which comes from the assumption of constant area for both solid and fluid phase) and no radial diffusion.

$$\frac{\partial C_f}{\partial t} + \frac{u}{\varepsilon} \frac{\partial C_f}{\partial z} = \frac{1 - \varepsilon}{\varepsilon} \frac{\partial C_s}{\partial t}$$
(70)

The Eq. 70 is equivalent to the equation presented by Reverchon [1].

## A.2.3. Momentum conservation

Similarly to the mass conservation, the momentum conservation is derived for inviscid fluid with no body forces

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}_f} \left( \rho_f V_z \right) d\mathcal{V}_f + \iint_{S} \left( \rho_f V_z \mathbf{V} \right) \mathbf{dS} = \iint_{S} (PdS)_z \tag{71}$$

where  $V_z$  is the z component of the velocity.

We the momentum conservation to the shaded control volume in Fig. 10, the integrals on the left side are evaluated in the same manner as discussed above in the regard to the continuity equation. That is,

$$\frac{\partial}{\partial t} \iiint_{\mathcal{V}_f} \left( \rho_f V_z \right) d\mathcal{V}_f = \frac{\partial}{\partial t} \left( \rho_f V A_f dz \right) \tag{72}$$

equation

and

$$\iint_{S} (\rho_{f} V_{z} \mathbf{V}) dS = -\rho_{f} V^{2} + (\rho_{f} + d\rho_{f}) (V + dV)^{2} (A + dA)$$
(73)

The evaluation of the pressure force term on the right side of Eq. 71 can be understood based on the Fig. 11. Here, the z components of the vector PdS are shown on all four side of the control volume. Remember that dS is

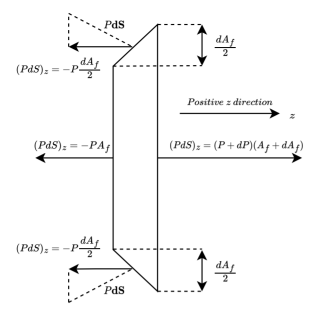


Figure 11: The forces in the z direction acting on the control volume

assumed to points away from the control volume; hence any z component  $(PdS)_z$  that acts toward the left (in the negative z direction) is a negative quantity, and any z component that acts toward the right (in the positive z direction) is a positive quantity. Also note that the z component of PdS acting on the top and the bottom inclined faces of the control volume in Fig. 11 can be expressed as the pressure P acting on the component of the inclined are projected perpendicular to the z direction,  $dA_f/2$ ; hence, the contribution of each inclined face (top or bottom) to the pressure integral in Eq. 71 is  $-P(dA_f/2)$ . All together, the right-hand side of Eq. 71 is expressed as follows:

$$\iint (PdS)_z = -PA_f + (P+dP)(A+dA_f) - 2P\frac{dA_f}{2}$$
 (74)

Substituting Eqs. 72 to 74 into Eq. 71, we have

$$\begin{split} &\frac{\partial}{\partial t} \left( \rho_f V A_f dz \right) - \rho_f V^2 A_f + (\rho_f + d\rho_f) (V + dV)^2 (A_f + dA_f) \\ &= P A_f - (P + dP) (A + dA_f) + P dA_f \end{split} \tag{75}$$

Cancelling like terms and ignoring products of differentials, equation above becomes in the limit dz becoming very small

$$\frac{\partial}{\partial t} \left( \rho_f V A_f dz \right) + d \left( \rho_f V^2 A_f \right) = -A dP \tag{76}$$

Dividing above equation by dz and taking the limit as dz goes to zero, we obtain

$$\frac{\partial \left(\rho_{f} V A_{f}\right)}{\partial t} + \frac{\partial \left(\rho_{f} V^{2} A_{f}\right)}{\partial z} = -A_{f} \frac{\partial P}{\partial z} \tag{77}$$

The Eq. 77 can be expanded further by assuming that  $A_f = \mathbf{A}\varepsilon$ 

$$\frac{\partial \left(\rho_{f}V\mathbf{A}\varepsilon\right)}{\partial t} + \frac{\partial \left(\rho_{f}V^{2}\mathbf{A}\varepsilon\right)}{\partial z} = -\mathbf{A}\varepsilon\frac{\partial P}{\partial t} \tag{78}$$

The equation can be further simplified by assuming that the cross-section of an extractor **A** is constant and cancel out

$$\frac{\partial \left(\rho_f V \varepsilon\right)}{\partial t} + \frac{\partial \left(\rho_f V^2 \varepsilon\right)}{\partial z} = -\varepsilon \frac{\partial P}{\partial t} \tag{79}$$

If the superficial velocity  $u = \varepsilon V$  is introduced, then the momentum conservation becomes

$$\frac{\partial \left(\rho_f u\right)}{\partial z} + \frac{\partial \left(\rho_f u^2 / \varepsilon\right)}{\partial z} = -\varepsilon \frac{\partial P}{\partial z} \tag{80}$$

Eq. 77 represents the conservative form of the momentum equation for the quasi-one-dimensional flow. The equivalent non-conservative form can be obtained by multiplying the continuity equation by V and subtracting it from Eq. 77

$$\frac{\partial \left(\rho_{f}VA_{f}\right)}{\partial t}-V\frac{\partial \left(\rho_{f}A_{f}\right)}{\partial t}+\frac{\left(\rho_{f}V^{2}A_{f}\right)}{\partial z}-V\frac{\left(\rho_{f}VA_{f}\right)}{\partial z}=-A_{f}\frac{\partial P}{\partial z} \eqno(81)$$

Expanding the derivatives on the left-hand size of above equation and cancelling like terms, gives

$$\rho_f A_f \frac{\partial V}{\partial t} + \rho_f A_f V \frac{\partial V}{\partial z} = -A_f \frac{\partial P}{\partial z}$$
 (82)

Dividing above equation by  $A_f$  the non-conservative form of the momentum can be obtained

$$\rho_f \frac{\partial V}{\partial t} + \rho_f V \frac{\partial V}{\partial z} = -\frac{\partial P}{\partial z} \tag{83}$$

The Eq. 83 is stylistically the same as the general momentum conservation for one-dimensional flow with nobody forces. The momentum equation can be expressed in terms of superficial velocity  $u = V\varepsilon$ .

$$\rho_f \frac{\partial \left( u/\varepsilon \right)}{\partial t} + \rho_f \frac{u}{\varepsilon} \frac{\partial \left( u/\varepsilon \right)}{\partial z} = -\frac{\partial P}{\partial z} \tag{84}$$

By expanding all the terms of equation above, we get

$$\frac{\rho_f}{\varepsilon} \frac{\partial u}{\partial t} + \rho_f u \frac{\partial \varepsilon^{-1}}{\partial t} + \rho_f \frac{u}{\varepsilon} \frac{1}{\varepsilon} \frac{\partial u}{\partial z} + \rho_f \frac{u}{\varepsilon} \frac{\partial \varepsilon^{-1}}{\partial z} = -\frac{\partial P}{\partial z}$$
 (85)

If the bed is not compressible and doesn't change its properties during the batch, then  $\frac{\partial \varepsilon}{\partial t} = 0$ 

$$\frac{\rho_f}{\varepsilon} \left( \frac{\partial u}{\partial t} + \frac{u}{\varepsilon} \frac{\partial u}{\partial z} + u^2 \frac{\partial \varepsilon^{-1}}{\partial z} \right) = -\frac{\partial P}{\partial z}$$
 (86)

If the porosity is constant along an extractor, then the momentum conservation equation becomes

$$\frac{\rho_f}{\varepsilon} \left( \frac{\partial u}{\partial t} + \frac{u}{\varepsilon} \frac{\partial u}{\partial z} \right) = -\frac{\partial P}{\partial z}$$
 (87)

The Eq. 87 represents non-conservative form of the momentum equation for quasi-one-dimensional flow with no body forces and constant porosity.

#### A.2.4. Energy conservation

Let's consider the integral form of the energy equation for adiabatic flow with no body forces and no viscous effects

$$\frac{\partial}{\partial} \iiint_{\mathcal{V}_f} \rho_f \left( e_f + \frac{V^2}{2} \right) d\mathcal{V}_f + \iint_{S} \rho_f \left( e_f + \frac{V^2}{2} \right) \mathbf{V} \cdot \mathbf{dS} = - \iint_{S} (P\mathbf{V}) \cdot \mathbf{dS}$$
(88)

Applied to the shaded control volume in Fig. 10, and keeping in mind the pressure forces shown in Fig. 11, Eq. 88 becomes

$$\frac{\partial}{\partial t} \left[ \rho_f \left( e_f + \frac{V^2}{2} \right) A_f dz \right] - \rho_f \left( e_f + \frac{V^2}{2} \right) V A_f$$

$$+ \left(\rho_f + d\rho_f\right) \left[e_f + de_f + \frac{(V + dV)^2}{2}\right] (V + fV) \left(A_f + dA_f\right)$$

$$= - \left[-PVA_f + (P + dP)(V + dV) \left(A_f + dA_f\right) - 2\left(PV\frac{dA_f}{2}\right)\right]$$
(89)

Neglecting products of differential and cancelling like terms, the above equation becomes

$$\frac{\partial}{\partial t}\left[\rho_{f}\left(e_{f}+\frac{V^{2}}{2}\right)A_{f}dz\right]+d\left(\rho_{f}r_{f}A_{f}\right)+\frac{\left(\rho_{f}V^{3}A_{f}\right)}{2}=-d\left(PA_{f}V\right)$$

$$\tag{90}$$

Of

$$\frac{\partial}{\partial t} \left[ \rho_f \left( e_f + \frac{V^2}{2} \right) A_f dz \right] + d \left[ \rho_f \left( e_f \frac{V^2}{2} \right) V A_f \right] = -d \left( P A_f V \right)$$
(91)

Taking the limit as dz approaches zero, the equation above becomes the following partial differential equation

$$\frac{\partial \left[\rho_f\left(e_f + V^2/2\right)A\right]}{\partial t} + \frac{\partial \rho_f\left(e_f + V^2/2\right)VA_f}{\partial z} = -\frac{\partial \left(PA_fV\right)}{\partial z} \tag{92}$$

Equation 92 is the conservation form of the energy expressed in terms of the total energy  $e + V^2/2$ , appropriate for unsteady, quasi-one-dimensional flow. The energy equation can be expressed in terms of internal energy if Eq. 77 is multiplicated by V and then subtracted from Eq. 92

$$\frac{\partial \left(\rho_f e_f A_f\right)}{\partial t} + \frac{\partial \left(\rho_f e_f V A_f\right)}{\partial z} = -P \frac{\partial A_f V}{\partial z} \tag{93}$$

The equation above is the conservation form of the energy equation expressed in terms of internal energy  $e_f$  suitable for quasi-one-dimensional flow. The non-conservative for is then obtained by multiplying the continuity equation 58, by  $e_f$  and substructing it from 93, yielding

$$\rho_f A_f \frac{\partial e_f}{\partial t} + \rho_f A_f V \frac{\partial e_f}{\partial z} = -P \frac{\partial (A_f V)}{\partial z} \tag{94}$$

Expanding right-hand side and dividing by  $A_f$ , the above equation becomes

$$\rho_f \frac{\partial e_f}{\partial t} + \rho_f V \frac{\partial e_f}{\partial z} = -P \frac{V}{A_f} \frac{\partial A_f}{\partial z}$$
 (95)

or

$$\rho_f \frac{\partial e_f}{\partial t} + \rho_f V \frac{\partial e_f}{\partial z} = -P \frac{\partial V}{\partial z} - PV \frac{\partial (\ln A_f)}{\partial z}$$
(96)

Equation 96 is the non-conservative for of the energy equation expressed in therms of internal energy, appropriate to unsteady quasi-one-dimensional flow. The reason for obtaining the energy equation in the form of Eq. 96 is that, for a calorically perfect gas, it leads directly to a form of the energy equation in therms of temperature T. For calorically perfect gas  $e_f = C_n T$ 

$$\frac{\partial \left( \rho_f e_f A_f \right)}{\partial t} + \frac{\partial \left( \rho_f e_f V A_f \right)}{\partial z} = -P \frac{\partial A_f V}{\partial z} + \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) \quad (97)$$

## A.3. Bayes theorem

As discussed by Himmelblau [21], the Bayesian approach to estimation make use of prior information. Such prior knowledge can come from theoretical considerations, from the results of previous experiments, or from assumptions by the experimenter. Typically, a Bayesian approach assumes a prior probability distribution of an unknown parameter  $\theta$  in some parameter space  $\theta$ . The distribution is updated by using Bayes' rule to obtain the posterior probability distribution.

Consider a set of events or outcomes,  $A_1, A_2, ..., A_n$ , and some other event B. Bayes' theorem states that the probability that event  $A_i$  will occur, given that event B has already occured, which will denoted by  $P\{A_i|B\}$ , is equal to the product of the probability that  $A_i$  will occur regardless of whether B will take place and the probability that B will occur, given that  $A_i$  has already taken place, divided by the probability of the occurrence of B:

$$P\{A_i|B\} = \frac{P\{B|A_i\}P\{A_i\}}{P\{B\}}$$
 (98)

Further, if all events comprising the set  $\{A_i\}$  are included in  $A_1, A_2, ..., A_n$ , then

$$P\{A_i|B\} = \frac{P\{B|A_i\}P\{A_i\}}{\sum_{i=1}^{n} P\{B|A_i\}P\{A_i\}}$$
(99)

We can interpret these symbols as follows:

- P{A<sub>i</sub>} is a measure of our degree of belief that event
  A<sub>i</sub> will occur or that hypothesis A<sub>i</sub> is true prior to the
  acquisition of additional evidence that may alter the
  measure. P{A<sub>i</sub>} is denoted the prior probability.
- 2.  $P\{A_i|B\}$  is a measure of our degree of belief that event  $A_i$  will occur or that hypothesis  $A_i$  is true, given additional evidence B pertinent to the hypothesis.  $P\{A_i|B\}$  is termed the *posterior probability*.
- 3.  $P\{B|A_i\}$  denotes the likelihood that event B will occur, given that event  $A_i$  is true.  $P\{B|A_i\}$  is a conditional probability, interpreted in the Bayesian framework as a likelihood,  $L(A_i|B)$ .

For continuous variable, Bayes' theorem can be more conveniently expressed in terms of the probability density function rather than the probabilities themselves. Equation 99 can be expressed in terms of a set of observed values of the random variable X, the experimental data  $\mathbf{x}$  and unknown parameter  $\theta$  as

$$p(\theta|X = \mathbf{x}) = p(\theta|\mathbf{x}) = \frac{L(\theta|\mathbf{x})p(\theta)}{\int_{-\infty}^{+\infty} L(\theta|\mathbf{x})p(\theta)d\theta}$$
(100)

where

 $p(\theta|\mathbf{x})$  = the posterior probability density function for  $\theta$ ; it includes knowledge of the possible values of  $\theta$  gained from the experimental data  $\mathbf{x}$ 

 $p(\theta)$  = the prior probability density function for  $\theta$  (before the experiment in which **x** was observed)

 $L(\theta|\mathbf{x}) = p(\mathbf{x}|\theta)$  = the probability density function termed the likelihood function of  $\theta$  given  $\mathbf{x}$ 

The denominator in Equation 100 is a normalizing factor chosen so that the integration of the posterior distribution is unit, i.e.,  $\int_{-\infty}^{+\infty} p(\theta|\mathbf{x}) = 1$ . By taking into account the law of total probability, the denominator can be written as

$$\int_{-\infty}^{+\infty} p(\mathbf{x}|\theta) p(\theta) d\theta = p(\mathbf{x})$$
 (101)

If the prior distribution is a uniform distribution, that is the prior distribution is a constant, the Equation 100 reduces to

$$p(\theta|\mathbf{x}) = \frac{L(\theta|\mathbf{x})}{\int_{-\infty}^{+\infty} L(\theta|\mathbf{x}) d\theta}$$
(102)

If the prior knowledge concerning a postulated event or hypothesis is poor, the posterior probability is largely or entirely determined by the likelihood function, that is, by the additional accumulated evidence for which the likelihood function acts as a mathematical expression. If prior knowledge outweighs recent evidence, however, then the posterior probability is determined almost solely by the prior probability.

In the application of tests and the design of experiments, certain definitions and rules concerning probability are needed and are listed below.

- 1. It follows from the frequency theory of probability that: 0 < P < 1
- If the probability of occurrence of one event A depends on whether or not event B has occurred, the two events are termed dependent; if the probability of occurrence of event A does not depend on the occurrence of B, or the reverse, the two events are independent.
- 3. **Addition Rule** If  $A_1, A_2, ..., A_n$  are mutually exclusive events, i.e., cannot occur at the same time, the probability of occurrence of just one of the events is equal to the sum of the probabilities of each  $A_i$ :

$$P(A_1 \text{ or } A_2 ... \text{ or } A_n) = \sum_{i=1}^n P(A_i)$$
 (103)

Very often we let

$$\sum_{i=1}^{n} P\left(A_i\right) = 1 \tag{104}$$

Also, if each event is equi-probable so that  $P(A_i) = q$ ,

$$\sum_{i=1}^{n} q = nq = 1 \qquad \text{or} \qquad q = \frac{1}{n} = P(A_i)$$
 (105)

In set theory, mutually exclusive event have no points in common. The union of the sets which represents the set of all elements that belong to

$$P\left(A_{1} \cup A_{2} \cup ... \cup A_{n}\right) = P(A_{1}) + P(A_{2}) + ... + P(A_{n}) \ (106)$$

4. **Multiplication Rule** If *A* and *B* are *independent* event

$$P(A \text{ and } B) = P(A)P(B) \tag{107}$$

In set theory the intersection of *A* and *B* is the set of all elements that belong to *A* and *B*:

$$P(A \cap B) = P(A)P(B) \tag{108}$$

If the A and B are dependet events,

$$P(A|B) = \frac{P(A \cap B)}{P(B)} \tag{109}$$

where the symbol P(A|B) means "probability of A given B". As a corollary,

$$P(A \cap B) = P(B)P(A|B) \tag{110a}$$

$$= P(A)P(B|A) \tag{110b}$$

Two kind of probabilities enter Equation 110a (or Equation 110b): the absolute probability of event B (or A) irrespective of whether or not A (or B) has occurred, and that the conditional probability of event A (or B) computed on the assumption that B (or B) has occurred. Equation 108 or 109 is special case of Equation 110a or 110b, because if the event are independent P(A|B) = P(A). For the case of many event, Equation 108 can be expanded to

$$P(A_1 \text{ and } A_2 \text{ and } \dots \text{ and } A_n) = P(A_1) \cdot P(A_2) \cdot \dots \cdot P(A_n)$$
 
$$= \prod_{i=1}^n P(A_i)$$
 (111)

5. Another useful relationship for event which are not mutually exclusive is

$$P(A) + P(B) - P(A \cap B) = P(A \cup B)$$
 (112)

## A.4. Solid density measurement



## QUANTACHROME CORPORATION Upyc 1200e V5.05 Analysis Report

Tue Oct 11 15:10:37 2022

User ID: OLIW

## Sample Parameters

Sample ID: T5
Weight: 55.5411 g
Description: coal
Comment: Powder

Analysis Parameters Cell Size - Large

V Added - Large: 80.8546 cc

V Cell: 149.7915 cc

Analysis Temperature: 27.9 C
Target Pressure: 131.0 kPa
Type of gas used: Helium
Equilibration Time: Auto
Flow Purge: 1.0 min.

Maximum Runs: 5

Number Of Runs Averaged: 5
Deviation Requested: 0.0050 %

Analysis Results

Deviation Achieved: 0.1135 %
Average Volume: 44.1994 cc
Volume Std. Dev.: 0.0542 cc
Average Density: 1.2566 g/cc
Density Std. Dev.: 0.0015 g/cc
Coefficient of Variation: 0.1227 %

Run Data

RUN VOLUME (cc) DENSITY (g/cc)

1 44.1311 1.2585
2 44.1423 1.2582
3 44.2163 1.2561
4 44.2709 1.2546
5 44.2365 1.2555

 $\textbf{Figure 12:} \ \, \textbf{The result of solid density measurement}$