Modeling and simulation of drying operations in PVC powder production line: Experimental and theoretical study of drying kinetics on particle scale

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Abstract

Keywords:
- PVC
- Convective drying
- Fluidization
- Drying kinetic model
- Pneumatic dryer model

An experimental method to determine drying kinetic at a particle scale and a pneumatic dryer model are presented. The particle scale kinetics were obtained by immersion of a fixed mass of wet PVC particles (cake) in a batch dense fluidized bed containing inert hot particles (glass bead). It appears clearly that the PVC drying is controlled by a competition between internal and external transfers. The drying kinetic was described by a shrinking core type model and integrated in a one-dimensional steady-state model simulating a pneumatic dryer. A two-phase continuum model was used to describe the steady-state flow of a diluted dispersed phase (wet PVC powder) and a continuous phase (humid air) through dryer. The model takes into account the convective heat, mass and momentum transfers. The numerical results are compared with industrial experimental data. The results show that the inlet temperature is the most important parameter in the operation.

1. Introduction

This study has been realized in the Chemical Engineering Laboratory in Toulouse in partnership with INEOS ChlorVinyls. In PVC powder production line, after the polymerization step, a suspension, composed of water and PVC particles, is obtained. Most of this suspension water is eliminated during a centrifugation step, leading to a wet porous powder, called cake (with a humidity between 0.2 and 0.35 kg of water/kg of dry PVC). The cake drying occurs essentially in a pneumatic dryer coupled with a fluidized bed dryer with a role of eliminating the residual humidity of PVC (less than 0.05 kg of water/kg of dry PVC). This operation, which consumes a lot of energy, between 800 and 1500 kJ/kg dry PVC, in accordance to the operating conditions and the PVC grade, represents 30% of the production cost. With the purpose of reducing drying cost, it is important to understand the different physical phenomena involved (mass transfer, heat transfer, hydrodynamic phenomena specific to the dryers). This study focuses firstly on the acquisition of the kinetics data in a batch dense fluidized bed, and secondly on the modeling of the dehydration of PVC particles in a pneumatic dryer.

The drying kinetic is obtained by immersing a small amount of PVC particles (mean diameter: 143 μm) in a hot fluidized bed filled with glass beads (mean diameter: 324 μm). These bigger particles' role is to suppress the agglomeration phenomena between PVC particles and to realize the drying in isothermal conditions. The experimental results show that the drying kinetic is strongly dependent on air temperature and humidity.

These experimental results were represented according to a kinetic model based on heat and mass exchanges between PVC and air (external transfers), and heat and mass diffusion inside the particle (internal transfers). That kinetic model is then included in a steady-state one-dimensional pneumatic dryer model, which simulates the industrial dryer.

2. Background

Porous media drying is a complex problem and still not well-understood despite of the numerous studies that can be found in the literature, due to the coupled exchanges (mass, heat, and momentum transfers) between the gas and the particles [1], depending on solid particles’ morphological parameters and water/solid affinity. As shown in Fig. 1, during the drying of a porous particle, two periods can be distinguished. The first period consists in the elimination of free water located at its surface. In this period the evaporation is controlled by external transfers which can be, in accordance to the dryer technology, e.g., conductive, convective or radiative. In our case (pneumatic and fluidized bed dryer), these transfers are convective and depend on the local relative velocity between air and particles, called slip velocity. These transfers depending on the drying technology have been widely studied, and the literature provides a lot of correlations, sometimes
contradictory, for its estimation ([2–4]). In the second period, the water located in the particles' pores is eliminated: the drying is controlled by a combination between internal and external transfers. The internal transfers depend on the solid properties and structure. The literature shows that many elementary phenomena may occur during drying. Such phenomena are complex and depend on the structure of the solid particle, the water–solid liaison, and the water composition. These elementary mechanisms, represented in Fig. 2 can be ([5,6]):

- capillary forces, depending on liquid wetting on the solid surface ($\gamma_{LV} \cdot \cos(\theta)$), temperature and particle pore size and distribution. These last properties affect the saturation water vapor tension at the solid/liquid interface and the internal liquid movement in the pores,
- vapor diffusion through the pores under the influence of temperature and water vapor partial pressure,
- solid diffusion, corresponding to bound water migration on the solid surface. This mechanism occurs in the case of a solid partially dissolved in water and with a strong liaison between water and solid (chemical interaction, hydrogen liaisons, ...).

Thus, the modeling of a wet particle drying by hot air has to consider: water evaporation at the particle surface during constant drying rate period, and the movement of an evaporation zone between the wet core and the dry crust of the particle. This zone progresses from the particle surface to its core. Several numerical simulations were realized on porous material ([7,8]). The thickness of that drying zone is due to the wetting properties of the liquid in the pores: if the wetting is perfect, the liquid can spread on the pores' internal surface and affect the thermal gradient [9].

In the present case, the wetting properties of PVC, which depends on the superficial tension ($\gamma_{LV}$) of the aqueous solution introduced during the polymerization step and the affinity between the solid particles and this solution, are represented by the contact angle ($\theta$). The low values of liquid superficial tension and high values of the contact angle (between 80° and 94.7° [10,11]), lead to suppose that the drying zone could be modeled by a surface of discontinuity.

Concerning the capillarity effect, the pore-size distribution of a PVC particle, presented in Fig. 3, shows that the minimal pore size is about 30 nm. The Kelvin law expresses water activity ($p_w/p°$) as a function of pore radius and shows that, for pore radius higher than 10 nm, the water activity is equal to 1. Hence the capillary effect does not influence the liquid/water equilibrium in this study. Concerning the solid and liquid diffusion, previous analysis on INEOS’s PVC shows that it is non soluble in water and that the PVC has a small affinity.
with water. For example, in an atmosphere with 75% of relative humidity, the equilibrium humidity of PVC is 2.4 g of water/kg of dry PVC (at 50°C). Hence the liquid and solid diffusion can also be neglected in this study, and the only internal transfer remaining is the vapor diffusion. This mechanism is influenced by temperature and water vapor partial pressure gradients.

The SEM picture of a PVC particle (Fig. 4) shows a typical pellet–grain structure [12], indeed each particle is a primary particle stack. With such a structure two pore families can be distinguished (Fig. 3):

- the space between the primary particles (pore size superior to 250 nm), which is macroporosity (the average size of these pores is 3335.6 nm). These pores are dried in the pneumatic dryer (93% of pore volume).
- the primary particles’ own porosity (pore size inferior to 250 nm), which is mesoporosity (the average size of these pores is 81.6 nm). These pores are dried in the fluidized bed dryer (7% of pore volume).

A shrinking-core type model is used by numerous authors ([13–16]) to represent the evaporation front evolution with particles having a pellet–grain structure. In this kind of model, the particle is divided into two zones: a wet core, which shrinks over time, and a dry crust, through which the evaporated water diffuses to the particle surface. The elementary mechanisms involved here are conductive heat transfer (from particle surface to its core) and water vapor diffusion through the pores. If the heat transfer leads the drying phenomena, the problem is analogous to the classical Stefan problem for droplet evaporation or freezing ([17,18]).

Concerning the pneumatic dryer, this technology is commonly used in order to eliminate free or barely-bound water. A pneumatic dryer is a continuous, convective dryer with a diluted solid transport. Used in order to eliminate free or barely-bound water. A pneumatic dryer is a continuous, convective dryer with a diluted solid transport. The dryer is equipped with several temperatures (±0.1°C) and pressure (±0.1 mbar) sensors and two capacitive hygrometers (temperature and humidity sensors). As shown in Fig. 4, a vacuum pump collects the wet air to measure its humidity. The measurement system, whose role is to ensure that the humidity measurements are realized at constant conditions (air flow rate and temperature), is equipped with: a filter to prevent particles to damage the sensor, a thermostatic bath to maintain a constant temperature and prevent condensation phenomena, the hygrometer, and a rotometer to maintain a constant flowrate. This hygrometer measures the air dew point with an accuracy of 0.1°C. This hygrometer measurement is controlled according to a second hygrometer (which measures the relative humidity with an accuracy of 0.1%) placed at the cyclone outlet. To avoid

### Table 1

<table>
<thead>
<tr>
<th>Properties</th>
<th>PVC</th>
<th>Glass beads</th>
</tr>
</thead>
<tbody>
<tr>
<td>(d_{50}) (µm)</td>
<td>144</td>
<td>324</td>
</tr>
<tr>
<td>(d_{10}) (µm)</td>
<td>104</td>
<td>233</td>
</tr>
<tr>
<td>(d_{02}) (µm)</td>
<td>199</td>
<td>450</td>
</tr>
<tr>
<td>(d_{12}) (µm)</td>
<td>139</td>
<td>314</td>
</tr>
<tr>
<td>(\rho) (kg/m³)</td>
<td>1175</td>
<td>2500</td>
</tr>
<tr>
<td>(\tau) (s)</td>
<td>–</td>
<td>3.015</td>
</tr>
<tr>
<td>(x) (K)</td>
<td>–</td>
<td>37.7</td>
</tr>
<tr>
<td>(\lambda) (W/(m·K))</td>
<td>0.17</td>
<td>–</td>
</tr>
<tr>
<td>(\sigma) (J/(kg·K))</td>
<td>1670</td>
<td>–</td>
</tr>
<tr>
<td>(U_{diff}) (cm/s)</td>
<td>1.2</td>
<td>8.8</td>
</tr>
<tr>
<td>(U_{K}) (cm/s)</td>
<td>65.3</td>
<td>242.5</td>
</tr>
</tbody>
</table>

The experiments were realized with a cake provided by INEOS Chlorvinsyls France (the properties are presented in Table 1). The particle size analysis realized on dry particles shows a Gaussian-type distribution with a mean particle diameter of 144 µm. The characteristic sizes presented in Table 1 are determined by laser granulometry (Malvern Mastersizer 2000–Sirocco 2000 ± ±0.1 µm). The particle density, porosity, tortuosity, and pore-size distribution (Fig. 3) were determined by mercury porosimeter on dry product (Micrometritics – Autopore IV ± ±0.1 nm). To determine the cake humidity, different samples were placed in a ventilated oven at 50°C during 24 h: the cake initial humidity varies between 0.195 and 0.205 kg of water/kg of dry PVC.

Preliminary experiments realized in a convective dryer have shown that the drying kinetic strongly depends on the external conditions (sample weight and thickness, air flow rate, ...). The study of the physical phenomena at a particle scale is then difficult to realize. Hence, just as several studies realized in our laboratory, ([26,27]) the drying kinetic is studied by immersing a small sample of wet PVC (about 3 g) in a hot fluidized bed containing 1.3 kg of glass beads with a mean particle diameter of 324 µm (see properties in Table 1).

The experimental pilot is represented in Fig. 5. The dryer is composed of a 10 cm diameter and 40 cm height glass column. The presence of an extended section on the top prevents the fine particles to be elutriated. The fluidization air is pre-heated in an electrical heater, and when it is necessary, mixed up with water vapor produces by an evaporator as shown in Fig. 4. A water flow rate is spread on an electrical heater. The water is instantaneously evaporated and carried into the warm air stream. The fluidization flow rates (velocities) are chosen in agreement with the characteristic velocities (minimal fluidization and terminal settling velocity) of PVC particles and glass beads. These values determined at ambient temperatures are presented in Table 1. The air velocity range chosen is from 25 to 40 cm/s, which corresponds to 3 and 4.5 times the minimal fluidization velocity of the glass beads.

The dryer is equipped with several temperatures (±0.1°C) and pressure (±0.1 mbar) sensors and two capacitive hygrometers (temperature and humidity sensors). As shown in Fig. 4, a vacuum pump collects the wet air to measure its humidity. The measurement system, whose role is to ensure that the humidity measurements are realized at constant conditions (air flow rate and temperature), is equipped with: a filter to prevent particles to damage the sensor, a thermostatic bath to maintain a constant temperature and prevent condensation phenomena, the hygrometer, and a rotometer to maintain a constant flowrate. This hygrometer measures the air dew point with an accuracy of 0.1°C. This hygrometer measurement is controlled according to a second hygrometer (which measures the relative humidity with an accuracy of 0.1%) placed at the cyclone outlet. To avoid
condensation near the sensor, the pipe is heated with heater cables. Both hygrometers are coupled with a data acquisition system, which permits to record the temporal evolution of air humidity. A previous study, realized on the hygrometers, showed that the response times of the sensors are not neglectable regarding the operation time. Indeed, the response time to a humidity step signal (i.e., transition from a dry air atmosphere to a wet air atmosphere) is about 40 s while the time of the experiments occurring in the fluidized bed is about 1 min (Fig. 6). This study was realized with both hygrometers by switching from a dry air atmosphere to a wet air atmosphere (room atmosphere), according to the three-way valves for the hygrometer placed in the humidity measurement system, and by removing suddenly the sensor from the pilot. The sensor response is modeled by the association of a plug-flow reactor (representing the lag during the first seconds of the experiments) and a batch reactor with a time constant of 9.5 s (Fig. 5). The experimental and numerical responses of the hygrometer to a humidity step are presented in Fig. 5. As this transfer function models correctly the hygrometer behavior, it is used for the kinetic result exploitation and modeling. So it is important to differentiate the experimental kinetic, given by the raw results of the sensor, and the inherent kinetic: the real drying kinetic in the fluidized bed.

The temperature range chosen is from 30 to 60 °C: if the bed temperature is higher than 70 °C the PVC surface becomes sticky and can cause an agglomeration in the bed. The influence of surrounding air relative humidity is studied from 1.2 to 63.1%.

4. Methodology

The temporal evolution of the outlet air relative humidity (RH), recorded during each experiment, permits to calculate at each instant $Y$, the absolute air humidity (in kg of water/kg of dry air) and so, $W_{\text{drying}}$, the drying flow rate (in g of water/s), knowing the dry air flow rate ($F_d$ in kg/s) going through the dryer:

$$Y = 0.622 \frac{RH \cdot p(T_\text{h})}{P - RH \cdot p(T_\text{h})}$$

(1)

$$p(T_\text{h}) = \exp\left(\frac{16.83 \cdot T_\text{h}}{228.74 + T_\text{h}} + 6.4075\right)$$

(2)

$$W_{\text{drying}} = F_d \cdot Y$$

(3)

with $T_\text{h}$ as the temperature measured by the hygrometer (in °C), $p(T_\text{h})$ as the saturated water vapor pressure at this temperature, and $P$ as the absolute pressure in the dryer.

By integrating the curve representing the temporal evolution of the drying flow rate, the amount of water evaporated during the time $t$ ($m_w$) and the total amount of water evaporated during the experiments
The calculated initial humidity corresponds to the initial humidity measured in the ventilated oven at 50°C during 24 h with an error inferior to 5%.

The temporal evolution of the normalized solid humidity ($X/X_0$), represented in Fig. 7, is used to determine characteristic times, $t_{75}$, $t_{50}$, $t_{25}$, and the drying time, $t_{dry}$, correspond to the time necessary for the normalized solid humidity to reach, respectively, 0.75, 0.50, 0.25, and 0.05.

5. Preliminary results

The cake (wet PVC) is a very cohesive powder, so the particles tend to agglomerate themselves. This phenomenon reduces external and internal transfers. Indeed the bigger the particles are, the lower the convective transfer is. In the same way, the internal transfer is reduced with particle agglomerates as the water vapor has a longer diffusion path. So the evolution of outlet air humidity during batch drying is evaluated with cake samples of different masses. The initial calculated humidity was compared to the real sample’s humidity. This study showed that the optimal sample mass is between 2 and 4 g of cake. Indeed with inferior mass the sensor precision is too low, and with higher mass the sample is too sticky to disperse and the results are not reproducible. To reduce this phenomenon, the sample is mixed up with glass beads at ambient temperatures before being introduced in the fluidized bed. This proceeding breaks the bounds between wet particles, and so improves the sample dispersion in the bed. Fig. 8 shows that the glass bead mass ratio has an influence until 25%. For higher values, the drying time stays constant. Hence, the following experiments are realized with a glass bead mass ratio of 30%.

In Fig. 8 the evolutions of the normalized solid humidity versus time for different experiments realized in the same conditions (Exps. 1, 2, & 3) are presented. This figure shows that our experimental system is reproducible and the variance of $t_{50}$ is less than 1 s.

6. Experimental results

Table 2 sums up the experimental data (bed temperature, inlet air absolute humidity, and air velocity) and characteristic times: $t_{25}$, $t_{50}$.

![Graph](image)
Moreover, the saturation air humidity (see Eq. (5)) is presented in this table.

\[
Y^* = 0.622 \frac{P^*(T_{bed})}{P^* - P^*(T_{bed})}.
\] (5)

The effect of bed temperature, air velocity, and air initial humidity on the drying kinetic is presented in Figs. 10, 11, and 12. They show that the drying is extremely fast even at low temperature. In the operating conditions used, the drying time stays below 85 s. It is important to note that this is overestimated because of the hygrometer response time. These results also show that even in a strongly stirred dryer, as the fluidized bed, the external transfer resistance is not neglectable. These resistances, controlled by the hydrodynamic parameters of the system, determine the energy and mass flow exchange between the dryer atmosphere and the PVC particles.

Indeed, the experimental results in Fig. 10 show that a rise of the air flowrate from 9.2 to 12.4 kg/h decreases the drying time from 53 to 36 s. The higher the air velocity is, the higher the relative velocity between the air and the particles is. Hence, the mass and heat transfers are increased and the drying time is shorter. Regarding the bed temperature effect, the results presented in Fig. 11 show that a rise from 35 to 55 °C strongly decreases the drying time from 40 to 24 s. Indeed, a bed temperature increase will improve the heat transfer between the fluidized bed and the particles but also the mass transfer driving force \((Y^* - Y)\).

Fig. 12 shows the influence of air humidity on the drying kinetic and drying characteristic times and presents the evolution of the normalized solid humidity versus time. These results show that a rise of the air absolute humidity from 16 to 25 g of water/kg of dry air increases the drying time from 55 to 84 s. This can be explained by the effect of the drying driving force: a rise of initial humidity results to an increase of \(Y^*\) and so there is a drying driving force reduction.

All these results show that the drying kinetic of PVC particles is directed by both external and internal humidity transfer.

### 7. Kinetic model

From our experimental results and knowing the properties of the PVC powder, we determined a kinetic law. This law is implemented in a model simulating our experimental setup: the kinetic model. In this section, we described the hypothesis and the equations of this model, and, in order to validate it, we compare its results to our experimental results.
7.1. Hypothesis and equations

Knowing the low affinity of water with the PVC surface, the drying inside the particle can be represented by the moving of an evaporation front from the surface to the core of the particle (Fig. 13). The shrinking core model principle is based on the following hypotheses:

- The particles are considered perfectly spherical and uniform.
- The evaporation occurs on the evaporation front.
- The fluidized bed is considered as a stirred tank.
- The wet core humidity and the dry crust humidity are respectively considered equal to the particle initial humidity \( (X_0) \) and null.
- The mass transfer is directed by the water vapor diffusion through the dry crust and the convection at the particle surface.
- The heat is transferred from the air to the particle by convection and is diffused in the dry crust by conduction.
- The sample weight is low enough to assure the bed isothermlicity.
- The particle temperature is considered uniform. This hypothesis can be justified by the low value of the thermal Biot number. In addition, this hypothesis has been justified with an analog model calculating the temperature profile in the dry crust.
- The instantaneous profile of the air humidity in the dry crust is obtained with the quasi-stationary state hypothesis.

The equations of the model, solved with the variable steps Runge–Kutta method, are presented below:

- Water vapor flux density at particle surface:
  \[
  N_{\text{wm}} = \frac{k_y}{B_{\text{M}} (1 - R/r_c)} + 1 \cdot (Y' - Y). 
  \]  
  \( N_{\text{wm}} \) (6)

- with \( B_{\text{M}} \) the mass Biot number, defined as:
  \[
  B_{\text{M}} = \frac{k_y \cdot R}{D_{\text{app}} \cdot \rho_g}. 
  \]  
  \( B_{\text{M}} \) (7)

- Mass balance on the particle:
  \[
  \frac{dX}{dt} = - \frac{6 \cdot N_{\text{wm}}}{\rho_p \cdot d_p}. 
  \]  
  \( \frac{dX}{dt} \) (8)

- Instantaneous solid humidity:
  \[
  X = \left( \frac{d}{d_p} \right)^3 \cdot X_0. 
  \]  
  \( X \) (9)

- Wet core radius evolution:
  \[
  \frac{dr_c}{dt} = \frac{-k_y \cdot R^2}{\rho_p \cdot r_c \cdot |B_{\text{M}}| (R - r_c) + r_c} \cdot \frac{Y' - Y}{X_0}. 
  \]  
  \( \frac{dr_c}{dt} \) (10)

- Heat balance on the particle
  \[
  \rho_p \cdot C_p \cdot \frac{dT_p}{dt} = \frac{3 \cdot h \cdot (T - T_p)}{R} \left( \frac{dX}{dt} \right) \cdot \rho_p \cdot \Delta H_v \cdot \frac{dX}{dt}. 
  \]  
  \( \frac{dT_p}{dt} \) (11)
with \( C_p_h \), the specific heat of the wet particle, defined as:

\[
C_p_h = C_p_p + X \cdot C_p_t
\]

To solve these equations, the apparent diffusion and the mass and heat transfer coefficients are calculated using the below correlations:

- The apparent diffusion coefficient is calculated from the binary molecular diffusion coefficient.

\[
D_{app} = D_{air-H_{2}O} \frac{X}{\tau}
\]

- with \( \chi \) and \( \tau \) given in Table 1. The binary molecular diffusion coefficient is calculated from the Fuller, Schettler, and Giddings correlation [28]:

\[
D_{air-H_{2}O} = 1.1757 \times 10^{-8} \cdot T_{g}^{1.75}
\]

- The heat transfer coefficient is calculated from the Baeyens correlation [29]:

\[
Nu = 0.15 \cdot Re
\]

- The mass transfer coefficient is calculated from the Chilton and Colburn analogy, which is classically used in drying:

\[
\frac{h}{k_y} = \frac{C_p_h}{(Sc \cdot Pr)^{2/3}}
\]

7.2. Model validation

In Figs. 14 and 15 typical numerical results are presented. These results show that the real drying time is about 11 s, which is four
times faster than the time observed with the hygrometer and that the drying operation occurs in three phases:

- The sample warming-up period, from 0 to 3 s (phase 1).
- The constant drying rate period, from 3 to 10.5 s (phase 2).
- The falling drying rate period, from 10.5 to 13 s (phase 3).

In order to compare the numerical results with the experimental data, the hygrometer transfer function is integrated in the model. The theoretical results are compared with the experimental finding (Exps. 1, 2, and 3). Fig. 15 indicates a good concordance between the model predictions and the experimental data.

To have a better understanding of the relative influence of the internal and external resistances to mass transfer, the sensitivity of the mass Biot number was studied. Its expression is remembered below:

\[ B_{IM} = \frac{k_v - R}{D_{app} \cdot \rho_g} = \frac{k_v/\rho_g}{D_{app}/R} \]  

\[ B_{IM} = \frac{\text{internal resistance}}{\text{external resistance}} \]  

In the previous simulation conditions, the Biot number estimated by the model is equal to 1.44, which means a competition between both transfer resistances. Fig. 16 shows the outlet air absolute humidity evolution versus time for different Biot number values (from 0.1 to 10 times the Biot number calculated by the model). These results show that the drying is longer and more difficult with the high Biot number. So in our case, the external resistance is not negligible in front of internal resistance. This phenomenon can be explained by the morphological parameters of PVC particles. The low internal resistance can be attributed to the evaporation of water inside the macroporosity of small particles and to the low affinity with water.

The model considers a monodisperse granulometry, while in reality the PSD shows a Gaussian-type distribution (\(d_{10}, d_{50}, \text{ and } d_{90}\) presented in Table 1). Fig. 17 shows that, in our case, the particle diameter has a small influence on the characteristic times. So, the monodisperse hypothesis does not alter significantly the model results.

Fig. 18 shows the comparison between the characteristic times given by the model and by the experimental data. These results show that the model predicts correctly the effect of: bed temperature (Fig. 17), inlet air humidity between 0 and 16 g of water/kg of dry air (Fig. 17), and fluidization air flow rate up to 11 kg/s (Fig. 17), on the characteristic times. Besides, at lower air fluidization air flow rate and at higher inlet air humidity (>16 g of water/kg of dry air) the model underestimates the characteristic times, especially \(t_{25}\).

This phenomenon can be explained by different reasons:

- The effect of inlet air humidity and fluidization air flow rate on the cake dispersion. Indeed, a rise of inlet humidity and a decrease of fluidization flowrate disfavor this phenomenon.
- The effect of air humidity on the local equilibrium, which is not taken into account.
- And by considering the PVC particle as a homogeneous media (i.e., a uniform pore size). Indeed, an increase of air humidity can lead to a slow down of the drying rate. Moreover, the presence of mesoporosity in the PVC particles can explain the gaps observed, especially at the
end of the drying operations. In that case, transfers are controlled by a combination between molecular and Knudsen diffusion.

8. Pneumatic dryer model

The aim of this study is to model a pneumatic dryer (macroscopic model) using the particle scale kinetics previously elaborated. As shown in Fig. 1, the drying occurs in two periods:

- the removal of surface water, which is governed by a convective mechanism.
- the removal of pore water, which is driven by convective and diffusive mechanisms. This period is modeled with the shrinking core model described in the previous parts.

8.1. Hypothesis and equations

In this model, a two-phase continuum model was used to describe the steady-state flow of a diluted dispersed phase (wet powder) and a continuous phase through a pneumatic dryer. The model is based on the following assumptions:

- Mass, energy and momentum balances occur between the two phases.
- The friction forces between the dispersed phase and the wall can be neglected.
- The continuous phase, composed of a mixture of water vapor and other gas, is considered as an ideal gas.
- The particles are spherical and composed of a homogeneous porous matrix.
- The particle size distribution is considered as monodisperse.
- The wall heat loss is neglected.

The mass, energy, and momentum balances developed in one-dimensional steady-state for the k-phase and the transport equations for solid and gas humidity, deducted from the mass balances are presented below. In this work, the k-phase can be either the gas phase \((k = g)\) or the particle phase \((k = p)\).

- Mass balance:
  \[
  \frac{d}{dz} (\alpha_k \cdot \rho_k \cdot U_k) = \Gamma_k
  \]
  with \(\Gamma_k\) the volume rate of mass transfer of the k-phase, expressed from the water vapor flow density \(N_w\):
  \[
  \Gamma_p = -\Gamma_g = -\alpha_p \cdot s_p \cdot N_w
  \]
  where \(\alpha_p\) is the local volumetric fraction of solid and \(s_p\) the specific surface area of a particle. The water vapor flux density can be expressed as the product of a "global mass-transfer coefficient" and the drying driving force:
  \[
  N_w = K_y \cdot (Y - Y). 
  \]
  - The global mass-transfer coefficient is expressed as a function of the mass Biot number (as seen in the previous part in Eq. (6)): 0.43
  \[
  K_y = \frac{k_y}{B_{IM}} \left( \frac{X}{X_C} \right)^{1/3} \left(1 - \frac{1}{X_0 - X_C} \right) 
  \]
  - Eq. (21) is valuable for solid humidity less than or equal to the critical humidity, \(X_C\):
  \[
  X_C = \frac{\rho_w \cdot X}{\rho_p}
  \]
  - This case is commonly observed when the pneumatic dryer is fed by a cake obtained by an efficient centrifugation. In opposite, when the initial humidity is greater than \(X_C\), the evaporation takes place essentially on a continuous liquid film at the particle surface (convective transfer). The reduction of this film thickness during the evaporation decreases the drying surface. In this case the global mass transfer coefficient can be estimated by:
  \[
  K_y = k_y \cdot \left( \frac{\rho_y \cdot X}{\rho_{water}} + 1 - X \right)^{2/3} \text{ for } X > X_C 
  \]
  - Energy balance:
  \[
  \int_0^L \frac{dH_k}{dz} = Q_{q\rightarrow k} + Q_{w\rightarrow k} + A_c \cdot \Gamma_k \cdot H_w
  \]
Humidity transport equations:

\[ Q_{g\rightarrow k} = -Q_{p\rightarrow g} = \alpha_p \cdot A_k \cdot s_p \cdot h \cdot (T_g - T_p) \]  \( \text{(25)} \)

where \( A_k \) is the pipe cross section, \( h \) is the heat-transfer coefficient, and \( T_g \) and \( T_p \) are the solid and gas phase temperatures, respectively. As the solid flow is very diluted, the heat loss of the solid phase is neglected; for the gas phase it is expressed as:

\[ Q_{w\rightarrow g} = h_{wall} \cdot (T_{wall} - T_g) \]  \( \text{(26)} \)

Momentum balance:

\[ \frac{d}{dz} \left( \alpha_k \cdot \rho_k \cdot U_k^2 \right) = \alpha_k \cdot \frac{dp}{dz} - \alpha_k \cdot \rho_k \cdot g \cdot l_{q\rightarrow k} + \left( U_p - U_k \right) \cdot I \cdot \frac{d}{dz} + F_{j, w\rightarrow k} \]  \( \text{(27)} \)

The first two terms on the right-hand side of this equation represent, respectively, the influence of pressure and gravity on the flows. \( l_{q\rightarrow k} \) represents the interactions between the phases and \( F_{j, w\rightarrow k} \) represents the friction forces between the wall and the k-phase. In this model, the only interaction taken into account is the drag:

\[ l_{q\rightarrow k} = -l_{p\rightarrow g} = \frac{3 \cdot \alpha_k \cdot \rho_k \cdot U_k^2 \cdot C_d}{4 \cdot D_p} \]  \( \text{(28)} \)

where \( U_r \) is the slip velocity \((U_r = U_k - U_p)\). The drag coefficient, \( C_d \), is calculated by the Wen and Yu correlation \([30]\):

\[ C_d = \frac{24}{Re} \left(1 + 0.15 \cdot Re^{0.687}\right) \cdot \alpha_k^{-1.7} \]  \( \text{(29)} \)

The friction force between the pipe and the continuous phase is estimated by:

\[ F_{j, w\rightarrow g} = \frac{f \cdot \rho_k}{2 \cdot D_{pipe}} \cdot U_k^2 \]  \( \text{(30)} \)

The friction factor, \( f \), is calculated by the Blasius formula:

\[ f = \frac{64}{Re_{pipe}} \quad \text{if} \quad Re_{pipe} \leq 2100 \]  \( \text{(31)} \)

\[ f = \frac{0.0791}{Re_{pipe}} \quad \text{if} \quad Re_{pipe} > 10,000. \]  \( \text{(32)} \)

As the solid flow is much diluted, the particle friction forces are neglected.

Humidity transport equations:

\[ \frac{dX}{dz} = \frac{A_k \cdot F_p}{F_p^g} \]  \( \text{(33)} \)

\[ \frac{dY}{dz} = \frac{A_k \cdot F_g}{F_g^k} = \frac{F_p}{F_p^g} \cdot \frac{dX}{dz} \]  \( \text{(34)} \)

• with \( Q_{p\rightarrow g} \) as the energy exchanged from the p-phase to the k-phase and \( Q_{w\rightarrow k} \) is the heat loss of the k-phase.

• with \( F_p^g \) and \( F_k^p \) as the dry flowrate of the solid and gas phases.

• Transfer coefficients

Some dimensionless empirical correlations can be used to calculate the heat-transfer coefficient in a pneumatic dryer:

- Baeyens correlation \([29]\) see Eq. (15).
- De Brandt correlation \([29]\):

\[ Nu = 0.16 \cdot Re^{1.2} \cdot Pr^{0.67} \]  \( \text{(35)} \)

- Bandrowski correlation \([31]\)

\[ Nu = 0.00114 \cdot \alpha_p^{-0.5984} \cdot Re^{0.8119} \]  \( \text{(36)} \)

The Chilton and Colburn analogy (see Eq. (16)) is used in order to calculate the mass transfer coefficient.

Baeyens and De Brandt correlations predict the transfer coefficient in a diluted phase. In order to model the effects of collisions between particles on the transfer rate, Bandrowski correlation \([31]\) expressed the Nusselt number as a function of the solid volume fraction, \( \alpha_k \). This relation indicates that in the inlet section of a pneumatic dryer, the important value of \( \alpha_k \) can counterbalance the high values of slip velocity. In other words, it takes into account the effect of solid dispersion in the inlet zone.

9. Results and discussion

Firstly, the model predictions are compared with the experimental measurements effectuated by Baeyens \([29]\) on an industrial pneumatic dryer. Secondly, they are confronted with our preliminary industrial results obtained at INEOS ChlorVinyls company.

The simulation data, PVC properties, geometrical and operating parameters, are resumed in Table 3.

9.1. Comparison with literature findings

In the case of the industrial experiments published by Baeyens \([29]\), the inlet PVC humidity (0.26) is greater than the critical humidity (0.134), so the drying rate is controlled by a two stage process. The first stage corresponds to the surface water evaporation and the second to the pore water evaporation. Fig. 19 represents the theoretical results obtained using the different correlations previously presented and the experimental measurements effectuated by Baeyens \([29]\) on an industrial process. Fig. 18 represents the evolution of solid humidity versus

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Simulation data ([29]).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data</td>
<td>Baeyens</td>
</tr>
<tr>
<td>Particle properties</td>
<td></td>
</tr>
<tr>
<td>Mean particles diameter ((\mu m))</td>
<td>180</td>
</tr>
<tr>
<td>Solid density (kg/m(^3))</td>
<td>1160</td>
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<tr>
<td>Inlet air temperature (C(^\circ))</td>
<td>127</td>
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<tr>
<td>Inlet air humidity (kg/kg)</td>
<td>0.003</td>
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<tr>
<td>Dry air flowrate (t/h)</td>
<td>46.48</td>
</tr>
<tr>
<td>Thermal loss (kW/m)</td>
<td>25</td>
</tr>
</tbody>
</table>

Operating parameters | | |
| Pipe diameter (m) | 1.25 | 1.7 |
| Pipe length (m) | 25 | 22 |
| Inlet air temperature (C\(^\circ\)) | 127 | 180 |
| Inlet air humidity (kg/kg) | 0.003 | 0.004 |
| Dry air flowrate (t/h) | 46.48 | 45 |
| Dry PVC flowrate (t/h) | 6.670 | 8.545 |
| Thermal loss (kW/m) | 25 | 25 |
while Fig. 18 represents the evolution of air temperature versus pipe length.

The differences observed between experimental data and theoretical results can be explained by the worst dispersion of the wet cohesive PVC in the gas phase at the pipe's inlet zone. The theoretical results which fit the best the experimental data were obtained using the Bandrowski correlation. This can be explained by the effect of the slip velocity on the Reynolds number, and so on the heat and mass transfer coefficients. As shown in Fig. 20, this magnitude is really important at the inlet and progressively decreases along the dryer. This leads to very important variations of the transfer coefficients (see Fig. 21). In the Bandrowski correlation, this effect is attenuated by the solid volume fraction of the dispersed phase.

9.2. Comparison with INEOS ChlorVinyls results

To establish the humidity and temperature profiles along the pneumatic dryer, INEOS’ industrial dryer is equipped with several hygrometers and thermocouples. Fig. 22 shows the comparison between industrial data and numerical results obtained using the Bandrowski correlation. It can be noticed that the temperature profile is correct while some differences appear on the drying flowrate evolution: in the industrial dryer, the humidity is barely constant in the first meters while in the model results, humidity rises strongly. This is due to the dispersion phenomenon (not taken into account in the model) which slows down the drying in the acceleration zone.

9.3. Parameter study

The results of the following simulations were obtained using the Bandrowski correlation and Baeyens parameters (see Table 3).

The effect of dry air mass flowrate was examined between 25 and 46.4 t/h. As shown in Fig. 23, this parameter has a small influence on the drying rate above 35 t/h. The air velocity influences the drying rate only in the acceleration zone, then the slip velocity is equal to the terminal settling velocity. At the dryer outlet the air is far to be saturated, so the decrease of the dry air flowrate does not affect the drying driving force significantly (e.g., a decrease of dry air flowrate from 46 t/h to 35 t/h increases the outlet solid humidity from 0.054 to 0.067 kg water/kg dry PVC, and a decrease of dry air flowrate from 35 t/h to 25 t/h increases the outlet solid humidity from 0.067 to 0.097 kg water/kg dry PVC).

Concerning the inlet air temperature, Fig. 24 shows a dominant effect (e.g., an increase of inlet air temperature from 96 °C to 156 °C decreases the outlet solid humidity from 0.097 to 0.014 kg water/kg dry PVC). Indeed, increasing the temperature leads to a more important heat-transfer driving force. But as the equilibrium humidity depends on the temperature, it will also increase the mass-transfer driving force.
The inlet air humidity effects were investigated between 0 and 0.0105 kg water/kg dry air. The last value corresponds to an atmospheric air (15 °C) almost saturated (98.4% of relative humidity). As shown in Fig. 25, such an increase of inlet air humidity increases the outlet solid humidity from 0.052 to 0.054 kg water/kg dry PVC. This increase shows that the inlet air humidity, which is a sustained parameter, slightly influences the drying driving force. It can be noticed that the solid temperature is influenced by:

- the dry air flow rate during the falling drying rate period only;
- the air humidity during the constant drying rate period only, and
- the air temperature influences it during both periods.

In fact the temperature stage value of the constant drying period is controlled by the drying driving forces. As shown in Eqs. (19) and (20), this force depends only on the air humidity, $Y_{fl}$, and the equilibrium humidity, $Y_e$, which is directly related to the solid temperature. In the falling drying rate period, the solid temperature increases until the thermal equilibrium with the air, this equilibrium is controlled by the dry air flow rate and temperature.

10. Conclusion

An experimental and numerical protocol was developed to study the drying kinetic of porous media at the particle scale. The results show the significant influence of the operating parameters, essentially air temperature and humidity. It appears clearly that the PVC drying is controlled by a competition between internal and external transfers. The use of a shrinking-core type model to simulate the particle-scale drying was confirmed by the experimental results.

PVC is essentially constituted of macropores ($d_{p,m} < 250$ nm). The volume fraction of mesopores ($10 < d_{p,m} < 250$ nm) is inferior to 10%. So we choose a shrinking core type model, supposing that the particles can be represented like a spheric pseudo-homogeneous sphere (i.e., uniform porosity and pore size). This model permits the drying kinetic to be simulated correctly. The gaps observed at the end of the drying operation is probably related to the transfers in the mesoporosity (primary particles).

A one-dimensional steady-state model for pneumatic dryer was established. This model is applied for the drying process of wet PVC powder. The drying rate is controlled by the convective transfer in the first period and by the convective and diffusive transfers in the second. The first period corresponds to the surface water drying; besides the second corresponds to the evaporation of the water in the pores, simulated by a shrinking core model. The model takes into account the convective heat, and mass and momentum transfers. The parameter study shows that the inlet temperature is the most important parameter in the operation. But this model does not take into account the wet powder dispersion, which is certainly the limiting step of this process.

### Nomenclature

<table>
<thead>
<tr>
<th>Roman symbols</th>
<th>Description</th>
<th>Unit</th>
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<tbody>
<tr>
<td>$A_f$</td>
<td>Pipe cross-section</td>
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</tr>
<tr>
<td>$C_d$</td>
<td>Drag coefficient</td>
<td>-</td>
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<tr>
<td>$C_p$</td>
<td>Specific heat</td>
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<tr>
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<td>Particle diameter</td>
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References


