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Experimental study of hydraulic transport of large particles in horizontal pipes

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Abstract

This article presents an experimental study of the hydraulic transport of very large solid particles (above 5 mm) in an horizontal pipe. Two specific masses are used for the solids. The solids are spheres that are large with respect to the diameter of the pipe (5, 10 and 15%) or real stones of arbitrary shapes but constant specific mass and a size distribution similar to the tested spherical beads. Finally, mixtures of size and / or specific mass are studied. The regimes are characterized with differential pressure measurements and visualizations. The results are compared to empirical models based on dimensionless numbers, together with 1D models that are based on mass and momentum balance. A model for the transport of large particles in vertical pipes is also proposed and tested on data available in the Literature, in order to compare the trends that are observed in the present experiments in a horizontal pipe to the trends predicted for a vertical pipe. The results show that the grain size and specific mass have a strong effect on the transition point between regimes with a stationary bed and dispersed flows. The pressure drops are moreover smaller for large particles in the horizontal part contrary to what occurs for vertical pipes, and to the predictions of the empirical correlations.

Keywords: Hydraulic transport, solid-liquid two-phase flow, bed friction, deep sea mining.

1. Introduction

The hydraulic transport of solid particles is a method widely used in chemical and mining industries. Many predictive models exist in the case of slurries transported under homogeneous regime, that is to say when the particle diameter is small compared to the flow length scales and the velocity of the carrier fluid is high compared to the settling velocity of a particle [1–4]. It is then possible to predict the pressure losses in horizontal or vertical pipes with sufficient accuracy. In recent years, the sharp increase in demand for raw materials makes it interesting exploitation of new resources, particularly the use of fields at the bottom of the ocean [5, 6]. In this case, the solids may be large with respect to the pipe diameter and the circuit would have complex shapes, including vertical parts, horizontal parts, and potentially bends and S-shapes in order to absorb the deformations caused by surface waves. For transport of large particles in vertical pipe, a predictive model based on the work of Newitt *et al.* [7] and Richardson *et al.* [8] is proposed and validated on a set of experimental data [9–11]. However, in horizontal, and a fortiori in geometries in S-shape, there are few models [1–4, 12–15] and the effects of specific mass and more specifically of very large particle size have not been systematically explored. One major difficulty in the case of transport of large particles and high specific mass comes from the various flow regimes that may be observed [1, 2, 4, 12, 13, 16, 17]:

when the speed of transportation increases, several transitions arise from regimes with a layer of solids at the bottom of the pipe that is at rest or that flows backwards in inclined pipes [13, 16, 17] to regimes with a moving bed and eventually to heterogeneous and pseudo-homogeneous suspensions at high mixture velocities.

The knowledge of the velocity above which the bed starts to move forward is of great interest with respect to operation of a production line. Below this limit the system may indeed plug. In the present study, experiments are carried out in order to better understand the effects of solid size and specific mass on this velocity and on the pressure drop.

The experimental set-up is presented in § 2. Some general considerations on the typical regimes and pressure drop curves that are specific to an horizontal pipe and an overview of few models are presented in § 3. The experimental results are presented and discussed in § 4. The main results that concern mono-disperse calibrated solid spheres of two different sizes and two different specific masses flowing in the horizontal pipe are presented in § 4.1. Mixture of spheres and rough stones of arbitrary shapes are also tested in § 4.2, in order to check to what extent the results obtained for spheres may represent an actual application. Several models are proposed and compared to the experiments in § 4.3. Conclusions and perspectives are then given in § 5.

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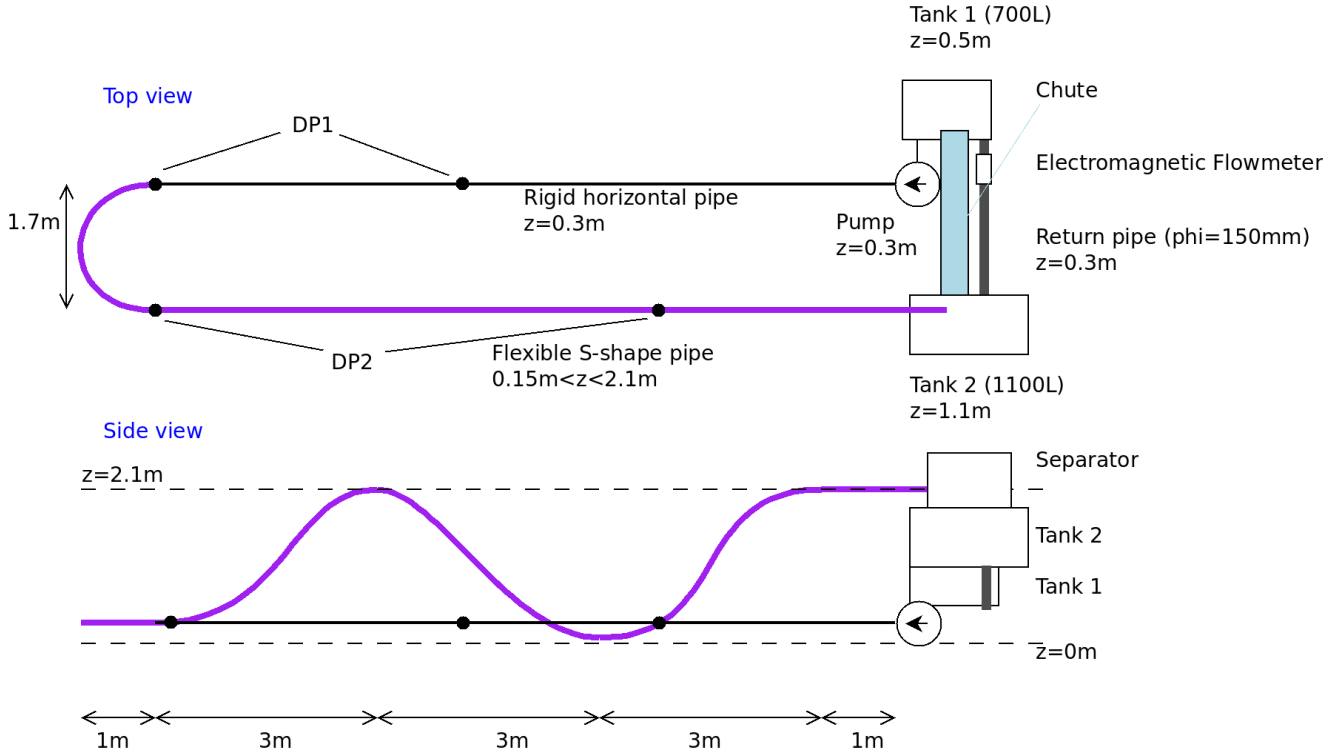


Figure 1: Sketch of the test loop. Top and side views.

2. Experimental set-up

2.1. Test loop

The test loop is shown schematically in Fig. 1. It consists of a first rigid, transparent horizontal pipe of internal diameter $D = 100$ mm and 10 m long in which flows the liquid/solid mixture. The return occurs in a clear flexible PVC hose reinforced with a steel coil, of internal diameter $D = 100$ mm and total length 20 m. This return pipe first follows a 180° horizontal curve of curvature diameter 1.7 m, then a climb, a descent and an ascent in a vertical plane. The present study focuses on the horizontal pipe.

The mixture of solids and water that flows in the rigid horizontal pipe and the return flexible pipe is then passed through a separator consisting of a closed box fitted with a side hatch and whose bottom side consists of a grid of stainless steel. The water then falls into the tank 2, and the solids flow in a chute that is inclined at 45° . This chute provides at its end an adjustable flap in order to control the solids flow rate. The end of the chute is above a grid container of known capacity that is immersed in tank 1 and connected with a grid pipe to the outlet of tank 1. The separated water flows from tank 2 into tank 1 through a pipe with an electromagnetic flow-meter. The mixture is sucked by a vortex pump (Ensival Moret MT 100-100-250) connected to the outlet of tank 1 and delivered in the circuit.

2.2. Characteristics of the solids

Solids with different sizes and specific masses are used. Their physical and geometrical characteristics are summarized in Tab. 1. The particles are relatively large, with sizes ranging from 5% to 18% of the pipe diameter. Calibrated beads of Glass (SiLi, SiLibeads type M, with a relative dispersion of sizes of 4%) and of Alumina (Umicore, Alumina Degussit 92%, with a relative dispersion of sizes of 10%) are used. The real solids that are used from the perspective of an actual application have irregular shapes, as can be seen in the picture in Tab. 1. The specific mass of a sample of fifty solids have been measured with a densimeter. It is constant within $2700 \pm 10 \text{ kg}\cdot\text{m}^{-3}$.

2.3. Control parameters and measured quantities

The aim of the present work is to measure the pressure drops in different parts of the test loop as a function of concentration and velocity. Several choices can be made for the definition of these quantities. The natural control parameters, *i.e.* the parameters that are really adjusted with experimental means, are the volumetric flow rates of the liquid (Q_l) and of the solids (Q_s).

The following set of parameters is chosen for presenting the results. The first is the mixture velocity V_{mix} and the second is the *transport* or *delivered* concentration C :

$$V_{mix} = \frac{Q_l + Q_s}{A}$$

$$C = \frac{Q_s}{Q_s + Q_l}$$







Type	Glass beads	Alumina beads	Mixture 1 (Alumina)	Mixture 2 (Alumina)	Mixture 3 (Alumina/Glass)	Stones
Size	5 mm 10 mm 16 mm	6 mm 15 mm	50% 6 mm 50% 15 mm D_{50} :10.5 mm	75% 6 mm 25% 15 mm D_{50} :8.25 mm	50% 6 mm Alumina 50% 5 mm Glass D_{50} :5.5 mm	8 – 18 mm D_{50} :10 mm
Specific mass	2500 kg.m ⁻³	3650 kg.m ⁻³	3650 kg.m ⁻³	3650 kg.m ⁻³	3075 kg.m ⁻³	2700 kg.m ⁻³
Picture						

Table 1: Physical characteristics of the calibrated beads and of the different mixtures. The mean diameter D_{50} is such that 50% of the solids are greater than this size.

with A the cross-section area of the pipe. This mixture velocity is a volumetric average of the velocities of each phase. It is convenient for comparison with models.

Concerning the concentration, please note that on the one hand, the solids do not flow with the same velocity as water, because of their large size and of the large density ratio —in other words there is a non-negligible *slip* velocity. On the other hand, in horizontal parts, among the various regimes that may be observed there exist regimes with a stationary layer of solids at rest at the bottom of the pipe [4, 12, 13]. There could thus be a great difference between the *transport* or *delivered* concentration C and the *in-situ*, *local* or *volumetric* concentration ϵ_s that is the ratio between the area occupied by the solids and the area of the pipe. Further, the volumetric concentration is a key parameter in the modelisation, as illustrated in § 3.1 and § 3.3. This concentration is however not a control parameter but results from the geometry, the physical properties of the phases and the boundary conditions.

The present experimental study is to perform measurements with varying V_{mix} in the range 0–5 m.s⁻¹, for constant delivered concentrations of 5, 10, 15 et 20%.

The water flow-rate is measured using an electromagnetic flowmeter (KROHNE Optiflux 2000) and adjusted by varying the rotation rate of the vortex pump. The solids flow-rate is set through a hatch and is measured by filling the buffer zone of known capacity located in the tank 1. Finally, the pressure drop are measured using two differential pressure sensors (VEGADIF65): a first that is located at the end of the straight line, 60 diameters downward of the pump, and a second that is located at the hose in the S-shaped part (see the position of the pressure taps in Fig. 1). After a stationary state is reached, the data are recorded for 30 s at a sample rate of 130 Hz. Measuring the rate of fluctuation of flow and pressure is used as a validation criterion of the measurements. The losses are expressed in terms of hydraulic gradients (meters of water column per meter of pipe) and with a dimensionless pressure drop coefficient:

$$I(m/m) = \frac{\Delta P}{\rho_l g L}$$

$$\lambda = \frac{\Delta P}{\frac{1}{2} \rho_l V_{mix}^2} \frac{D}{L}$$

with ΔP the measured static pressure drop, ρ_l the specific mass of the carrying liquid (water) and L the curvilinear distance between the pressure taps. The velocity scale that is used to present dimensionless results is the terminal velocity of the particle falling in the fluid at rest V_0 . Finally, the relative specific mass of the solids with respect to water is $s = \frac{\rho_s}{\rho_l}$.

In order to compare the present data to existing theories and correlations [4, 15], the following quantity is also introduced:

$$\Phi = \frac{I - I_f}{I_f}$$

with I_f the hydraulic gradient that would be observed for water flowing alone at the mixture velocity. It corresponds to the dimensionless excess of pressure drop caused by the presence of the particles. Similarly, two dimensionless groups are introduced for the mixture velocity:

$$F_D = V_{mix} / \sqrt{gD}$$

a “Froude” number based on the mixture velocity and the pipe diameter and

$$F_{dp} = V_0 / \sqrt{gd_p}$$

a “Froude” number based on the terminal velocity and the size of the particle.

In the following, I_h stands for the hydraulic gradient in the horizontal line, and I_s stands for the hydraulic gradient in the S-shape pipe. The case of the S-shape part is not fully developed here and will be discussed in a forthcoming paper with new experimental data including a better discretisation of the different inclined parts and bends. The symbol I_v is used for the hydraulic gradient that would be observed in a vertical pipe.

Optical measurements are also performed with a high-speed camera (Optronis CamRecord600). Typically 3200 images are recorded with a resolution of 1280×1024 pixels at a frame rate of 200 Hz. The flow is illuminated backwards with a LED plate from Phlox. The visualization area is surrounded by a square plexiglas box filled with water in order to minimize optical distortions.

3. Observed regimes and overview of a few models

A common feature of multiphase flows is the existence of very different flow patterns, which makes modeling much more complicated. In the case of the transport of large solid particles by a carrying liquid, several intermittent, stratified and dispersed regimes are observed in horizontal and inclined flows [1, 2, 4, 17]. For the specific case of vertical ascending flows, though plug or slug flows are observed in the pneumatic transport of small particles [18], to the best of our knowledge only dispersed regimes are observed for the hydraulic transport of large ($d_p \geq 1$ mm) and massive ($s \geq 1.5$) particles [2, 9–11]. The present section first presents a model for vertical ascending flows and then the typical regimes that are encountered in the horizontal experiment and the different possible models.

3.1. Validation of a model for the hydraulic transport of large particles in ascending vertical pipelines

The case of vertical flow is reasonably straightforward. The pressure force exerted on a column of fluid of height z balances two forces: the hydrostatic weight of the mixture and the friction on pipe wall due to the fluid shear stress [1, 2]. In the following, the hydrostatic weight of the column of water is removed in order to present the hydraulic gradients that are due to the flow of a mixture in the pipe.

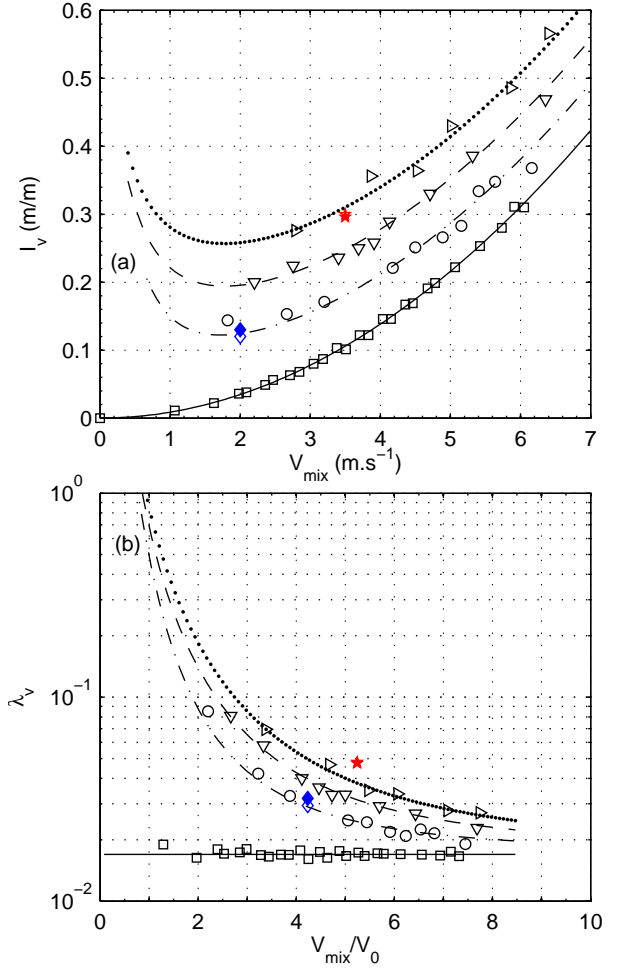


Figure 2: Validation of the vertical model on different data sets. Vertical hydraulic gradient I_v vs. mixture velocity V_{mix} . Black symbols: data of Yoon *et al.* [11] for spherical beads of diameter $d_p = 20$ mm, $s = 2.15$ flowing in a pipe of diameter $D = 100$ mm, compared to the results of the model (black lines) for various concentrations (\square and filled line: $C = 0\%$, \circ and dash-dotted line: $C = 5\%$, ∇ and dashed line: $C = 10\%$ and \triangleright and dotted line: $C = 15\%$). Red \star : data (open symbol) of Xia *et al.* [9] for spherical beads of diameter $d_p = 15$ mm, $s = 2$ flowing in a pipe of diameter $D = 100$ mm, at $C = 15\%$, compared to the result of the model (closed symbol). Blue \diamond : data (open symbol) of Hong *et al.* [10] for spherical beads of diameter $d_p = 5$ mm, $s = 2.5$ flowing in a pipe of diameter $D = 50$ mm, at $C = 3.85\%$, compared to the result of the model (closed symbol).

The vertical hydraulic gradient I_v can thus be decomposed into two parts: $I_v = I_{stat} + I_f$, with I_{stat} the hydrostatic contribution and I_f the wall shear-stress contribution. The hydrostatic contribution reads:

$$I_{stat} = (s - 1)\epsilon_s$$

The *in-situ* concentration ϵ_s is *a priori* unknown. Following the seminal work of Newitt *et al.* [7], the average velocity difference between the solids and the surrounding water, or *slip* velocity V_{slip} reads:

$$V_{slip} = \frac{1 - C}{1 - \epsilon_s} V_{mix} - \frac{C}{\epsilon_s} V_{mix} \quad (1)$$

This slip velocity would be the terminal velocity V_0 for a single solid of diameter d_p and of drag coefficient $c_d = 0.44$ falling in an infinite medium of fluid at rest:

$$V_0 = \sqrt{\frac{4d_p g(s-1)}{3c_d}} \quad (2)$$

However, it must be corrected in the case of a concentrated mixture flowing in a pipe. Owing to the range of parameters that we are interested in, we have chosen to use the Richardson & Zaki correlation [8] for the *hindered* average slip velocity:

$$V_{slip} = (1 - \epsilon_s)^{2.4} V_0 \quad (3)$$

The *in-situ* concentration is obtained by solving the non-linear system of Eqs. 1 and 3 in an iterative way, with a trust-region dogleg algorithm.

The model for the wall shear-stress contribution I_f is based on some assumptions. As noticed by Engelmann [5] or Hong *et al.* [10], large particles tend to migrate away from the wall due to hydrodynamic lift [2]. Assuming that the near-wall velocity profile is only slightly affected by the presence of particles in the core region, the wall shear-stress is modeled by water flowing at the water velocity:

$$I_f = \lambda \frac{(V_{mix} \frac{1-C}{1-\epsilon_s})^2}{2gD}$$

This model for vertical ascending flow has been validated on various experimental data available in the Literature [9–11]. The comparison between the model and the data is plotted in Fig. 2. The agreement is very good. When dealing with a mixture of liquid and solids, the pressure drops are significantly higher than for pure fluid for the whole range of mixture velocity that corresponds here to $V_0 \leq V_{mix} \leq 8V_0$: it is for instance twice as large at $C = 5\%$ and $V_{mix} = 4V_0$ for the beads of diameter $d_p = 0.2D$ and $s = 2.15$ (data of Yoon *et al.* [11], \circ in Fig. 2). A remarkable feature of the curves is the presence of a local minimum: the hydraulic gradient does not vary monotonically with the velocity. This is due to the fact that the hydrostatic gradient is the dominant term and that it is a decreasing function of the mixture velocity. In the case of ascending flows, the *in situ* concentration ϵ_s is indeed much larger than the *delivered* concentration C for mixture velocities of the order of V_0 and decreases slowly with increasing V_{mix} . For instance, for the data plotted with \circ in Fig. 2, $V_0 = 0.83 \text{ m.s}^{-1}$ and the values of the computed *in situ* concentration are: $\epsilon_s = 13\%$ for $V_{mix} = V_0$, $\epsilon_s = 11\%$ for $V_{mix} = 1.2V_0 = 1 \text{ m.s}^{-1}$, $\epsilon_s = 10\% = 2C$ for $V_{mix} = 1.4V_0$ and $\epsilon_s = 6\%$ for $V_{mix} = 4V_0$. The contribution of the hydrostatic gradient to the total hydraulic gradient is respectively 96%, 93%, 90% and 43%. The velocity which corresponds to the minimum of the pressure drop curve ($V_{mix} \simeq 1.8V_0$ for \circ in Fig. 2) is of great practical importance and the line should not be operated below this velocity.

The model is thus used with the parameters of the present experiments to compare the order of magnitude of hydraulic gradients and critical velocities between horizontal and vertical flows (see for instance Figs. 3 and 4b).

3.2. Description of the typical regimes in horizontal pipes

The Figure 3 presents a typical evolution of the hydraulic gradients I_h and I_s in the present experimental setup. The solids are Glass beads of diameter 5 mm and the delivered concentration is $C = 5\%$.

The thick black line in Fig. 3 stands for the measured hydraulic gradient in the horizontal part in the case with water flowing alone. The typical Reynolds number is 2×10^5 and the flow is fully turbulent. The curve is a fit of the form $I_f = \lambda \frac{V^2}{2gD}$ that gives a value for the friction coefficients $\lambda_h = 0.156$ and $\lambda_s = 0.160$. Corresponding rugosities can be deduced with the Colebrook formula. The estimated rugosity is $20 \mu\text{m}$ for the horizontal pipe.

The shape of the pressure drop curve with particles (red \bullet : I_h and blue \blacksquare : I_s in Fig. 3) is very similar to the ascending vertical case with a large increase of the pressure drop for the whole range of tested mixture velocity (here, $V_{mix} \leq 10.5V_0$). The presence of a local minimum is also observed: the hydraulic gradient does not vary monotonically with the velocity. The mixture velocity for which the minimum hydraulic gradient is observed will be termed the *critical velocity* V_{crit} , following the definition of Doron *et al.* [12]. In the present case and in the horizontal pipe, the critical velocity is $V_{crit,h} \simeq 1.8 \text{ m.s}^{-1}$ (*i.e.* $V_{crit,h} \simeq 3.8V_0$). All these observations are consistent with all the previous studies that dealt with similar large particles [4, 12, 14, 15].

For $V_{mix} < V_{crit}$ in the horizontal pipes, flow regimes with a stationary bed above which a compact layer of beads is flowing are observed (see bottom left picture in Fig. 3 that corresponds to $V_{mix} = 1.2 \text{ m.s}^{-1}$, *i.e.* $V_{mix} = 2.5V_0$). The more the mixture velocity decreases, the more the solids tend to settle down. These regimes are thus such that $\epsilon_s \gg C$ ($\epsilon_s \simeq 32\%$ for bottom left picture in Fig. 3) and are characterized by a large pressure drop that is caused by a decrease in the discharge section. In the ascending part of the flexible pipe, a layer of solids located at the bottom of the pipe and that is flowing backwards is even observed for these low velocities, as already observed by Yamaguchi *et al.* [17] (see top left picture in Fig. 3 that is taken at $V_{mix} = 0.9 \text{ m.s}^{-1}$). The flow in this regime is very unstable and the transit time needed to reach a stationary state is very long, of the order of twenty minutes—the typical time for a solid to flow through the whole pipe being 30 s.

Slightly above the critical velocity—for $V_{mix} \gtrsim V_{crit}$ —a moving bed on the bottom of the pipe is observed both in the horizontal pipe and in the S-shaped part (see bottom central picture and top right picture in Fig. 3 that are both taken at $V_{mix} = 2.1 \text{ m.s}^{-1}$, *i.e.* $V_{mix} = 4.1V_0$). Very few beads are also transported by saltation. The mean

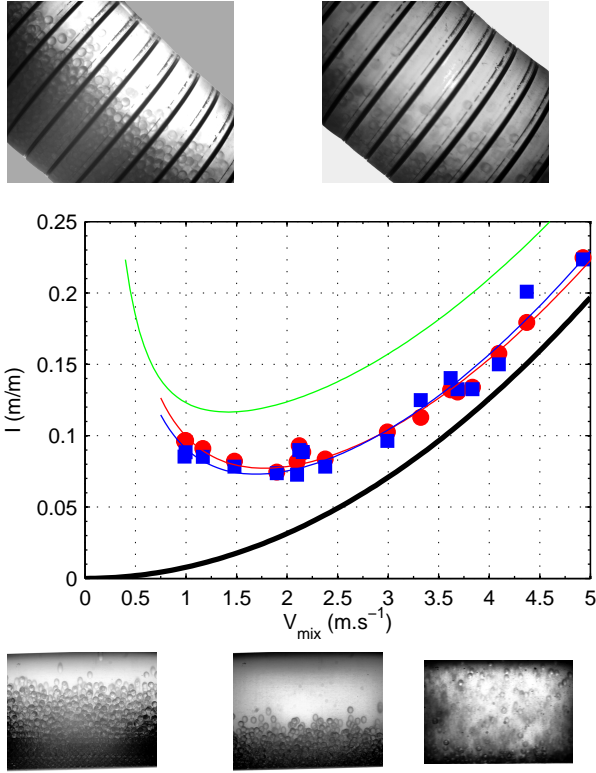


Figure 3: Illustration of the different regimes. Hydraulic gradient *vs.* V_{mix} , Glass beads of 5 mm, $C = 5\%$. Solid black line: water flowing alone I_f , red \bullet : I_h , blue \blacksquare : I_s . The blue and red solid lines are fits of the type $a/V_{mix} + bV_{mix}^2$ (see § 3.3). The green solid line stands for I_v predicted with the model developed for vertical pipelines (see § 3.1).

height of this moving bed is roughly 25 mm in the present case (Glass beads, $d_p = 5$ mm and $C = 5\%$), that corresponds to approximately five layers of beads. There is a velocity gradient between the upper layer of the moving bed and the lower layer that moves more slowly, as usually observed for hydraulic transport of sediment in two-dimensional open channels [19]. Please note that there may be substantial differences between the present experiments and usual bed-load models for the transport of sediments [19], because of three-dimensional and geometrical effects due to the cylindrical section of the pipe and of the large particle diameter to pipe diameter ratio. The average velocity of this bed is small compared to the mixture velocity and $\epsilon_s > C$ here, $\epsilon_s \simeq 10\%$.

Increasing further the mixture velocity, more and more solid beads get suspended and transported by the flow, there is no more bed, the flow is fully dispersed. The pressure drop curves behave as the clear-water pressure drop curve and follow the same trend at high velocities. In that case, $\epsilon_s \gtrsim C$ and the regime is called “pseudo-homogeneous” [12] or “heterogeneous” [4] (see bottom right picture in Fig. 3 that is taken at $V_{mix} = 4.9$ m.s⁻¹, *i.e.* $V_{mix} = 10.3V_0$).

The green line is the results of the model for pressure drop in ascending vertical flow, presented in § 3.1. The

main contribution to the increase of the total hydraulic gradient in the vertical model comes from the hydrostatic pressure of the mixture. On the contrary, in the case of the horizontal pipe, the increase in the total hydraulic gradient originates in the bed formation that causes more friction. Some partial conclusions concerning the comparison of a vertical ascending pipe and an horizontal pipe in view of an application with complicated shapes can be drawn. Firstly, the hydraulic gradient in the horizontal part is lower than the one that would be observed in an ascending vertical pipe at least for this specific mass and solid size. Secondly, the critical velocity in the horizontal pipe is greater than that of a vertical pipe. The effects of size, concentration and specific mass that may be very different for a vertical and an horizontal pipe will be further explored in § 4.

3.3. Correlations and models

This paragraph is a brief overview of some of the correlations and models that are commonly used.

Empirical correlations. The first quantity of interest to be presented is the critical velocity V_{crit} . One correlation has been proposed by Durand & Condolios (1952) [14]:

$$V_{crit} = F_l \{2Dg(s-1)\}^{1/2} \quad (4)$$

with F_l a constant of order unity, that depends on the delivered concentration and the particle size.

Concerning the prediction of the hydraulic gradient, a first empirical correlation is the one proposed by Durand & coworkers [15]. They used sand particles of one specific mass with diameter up to 25.4 mm in pipes ranging from 38 mm to 558 mm in diameter (the maximum relative diameter in the pipe of 104 mm was 4.5%). The dimensionless excess of pressure drop caused by the presence of the particles Φ is expressed using the two “Froude” numbers F_D and F_{d_p} . These two parameters are grouped to form $\Psi = F_D^2 F_{d_p}^{-1}$. They found a general correlation that best represents all their data:

$$\Phi = 180 C (\Psi)^{-1.5} \quad (5)$$

It is recommended to use this correlation in the vicinity of the critical velocity (from mixture velocities slightly below to three or four times greater). This correlation was then modified to take into account the specific mass of the particles, as reported by Newitt *et al.* [4]. In the case of particles of a few millimeters flowing in water, the particle Reynolds number is sufficiently high to assume that the settling velocity is given by Eq. 2 with a constant drag coefficient c_d . This form of the correlation reads:

$$\Phi = 121 C \left(\frac{V_{mix}^2}{gD(s-1)} \sqrt{\frac{3}{4}c_d} \right)^{-1.5}$$

The functional form of I_h that is predicted by this model is thus: $I_h = I_f (1 + AV_{mix}^{-3}) = bV_{mix}^2 + aV_{mix}^{-1}$, as displayed in Figs. 3 and 6.

This correlation have been widely discussed [1, 2, 4, 12]. The main criticisms are that it does not take into account the size of the particles and that a single correlation may not be applicable to all flow regimes. The value of the constant that is reported in the Literature is moreover different according to various authors, the term $F_{d_p}^{-1}$ being sometimes abruptly replaced by $\sqrt{c_d}$.

Semi-empirical correlations. Newitt *et al.* [4] have lead experiments in a 25.4 mm in diameter horizontal pipe, with various particles covering four specific masses ($s = 1.18$, $s = 1.4$, $s = 2.6$ and $s = 4.1$) and various sizes, the largest being of equivalent diameter 2.6 mm (10% of the pipe diameter). Theoretical considerations are used to derive expressions of the hydraulic gradient for different flow regimes (homogeneous, heterogeneous and flow with moving beds), and of the transition velocities between those regimes. For heterogeneous flows, *i.e.* for $17V_0 \leq V_{mix} \leq \sqrt[3]{1800gDV_0}$, the excess of friction is considered proportionnal to the energy dissipated by the particles as they fall under the action of gravity. This hypothesis leads to:

$$\frac{\Phi}{\epsilon_s} \propto (s - 1)V_0V_{mix}^{-3} \quad (6)$$

For flows with a moving bed, *i.e.* for $\sqrt{2gD(s - 1)} \lesssim V_{mix} \leq 17V_0$, the effect of the solids is proportional to the solid-solid friction between the solids and the bottom of the pipe:

$$\frac{\Phi}{\epsilon_s} \propto (s - 1)V_{mix}^{-2} \quad (7)$$

The main asumption in Ref. [4] is the use of C for ϵ_s , that is equivalent to assume that the particles move at the same speed as the water or at a constant fraction of it. The suspension of particles through turbulence or Bagnold forces is moreover not taken into account. This is obviously not the case in the present experiment as can be seen with the pictures in Fig. 3 and the values of ϵ_s/C that are reported in § 3.2.

Analytical model based on mass and momentum balance. Doron *et al.* [12] have established such a model. It is based on the decomposition of the cross-section of the pipe into two layers. It is thus a one dimensional model. The bottom of the pipe is assumed to be filled with a stationary or moving bed of packed particles. The height of this bed is y_b and the volumetric concentration in this layer is $C_b = 0.52$. All the particles are assumed to move at the same velocity in the bed. An heterogeneous mixture of solids and fluid is flowing in the upper part of the pipe. The mixture is treated as an homogeneous fluid with averaged physical properties and no slip between the phases is considered. The mass and momentum balance are then written in each layer. The shear stresses at the walls and at the interface between the two layers are modeled with frictions coefficients, and with a static friction force for the lower layer. In addition, the dispersion process of the

solid particles in the upper layer is modeled by a turbulent diffusion process balanced by the gravitational settling of particles, leading to an advection-diffusion equation. The size of the particles is taken into account, firstly to define the roughness of the interface between the two layers, and secondly in the definition of the turbulent diffusion coefficient and the advection velocity that is the hindered terminal velocity (the Richardson & Zaki correlation [8], Eq. 3, is used). This model leads to a non-linear system of 5 equations with 5 unknowns: the bed height (y_b), the velocity of the upper layer (U_h), the velocity of the lower layer (U_b), the concentration in the upper layer (C_h) and the pressure gradient (∇P). The parameters of the model are: the solid friction coefficient between the pipe wall and the particles (η), an angle of internal friction that models the normal stress transmitted by the shear stress at the interface between the fluid and the bed (φ), the packing concentration (C_b) and the correlations for fluid friction coefficients. This model have been implemented in Matlab, using an iterative procedure with a trust-region dogleg algorithm to solve the non-linear system. This implementation has been validated on the data of Ref. [12] in Fig. 4f.

4. Results and discussion

4.1. Effects of the physical characteristics of the beads

This paragraph is devoted to the comparison of the pressure drop curves with various concentrations, specific masses and sizes for identical spherical beads. The reference case is the Alumina beads of diameter 6 mm, of specific mass $s = 3.65$, and at a delivered concentration $C = 5\%$. In this case, the order of magnitude of the minimal pressure drop at critical velocity $V_{crit,h} \simeq 2.4 \text{ m.s}^{-1}$ is $I_{crit,h} \simeq 0.11 \text{ m/m}$.

The effects of the concentration are presented in Fig. 4a-b. Only results for the horizontal part are plotted. On the one hand, increasing the delivered concentration leads to an increase of the pressure drop, as expected. The relative increase of the pressure drop at the critical point with respect to $C = 5\%$ is roughly 40% for $C = 10\%$ and 70% for $C = 15\%$. Very few points are available for the concentration $C = 20\%$ owing to the large power required; these points are nevertheless on both sides of the critical point. The relative increase in pressure drop at the critical point for $C = 20\%$ is around 100%. On the other hand, changing the concentration seems to increase only very slightly the critical velocity (see § 4.3 for a discussion of this point).

The comparison of two sizes of beads of same specific mass at the same concentration is presented in Fig. 4c-d: the experimental measurements for Alumina beads of diameter $d_p = 6 \text{ mm}$ are plotted with \circ and the results for Alumina beads of diameter $d_p = 15 \text{ mm}$ are plotted with \square . The pressure drop that would be observed in an ascending vertical pipe according to the prediction of the model (see § 3.1) is plotted with, respectively a solid red line for $d_p = 6 \text{ mm}$ and a dashed red line for $d_p =$

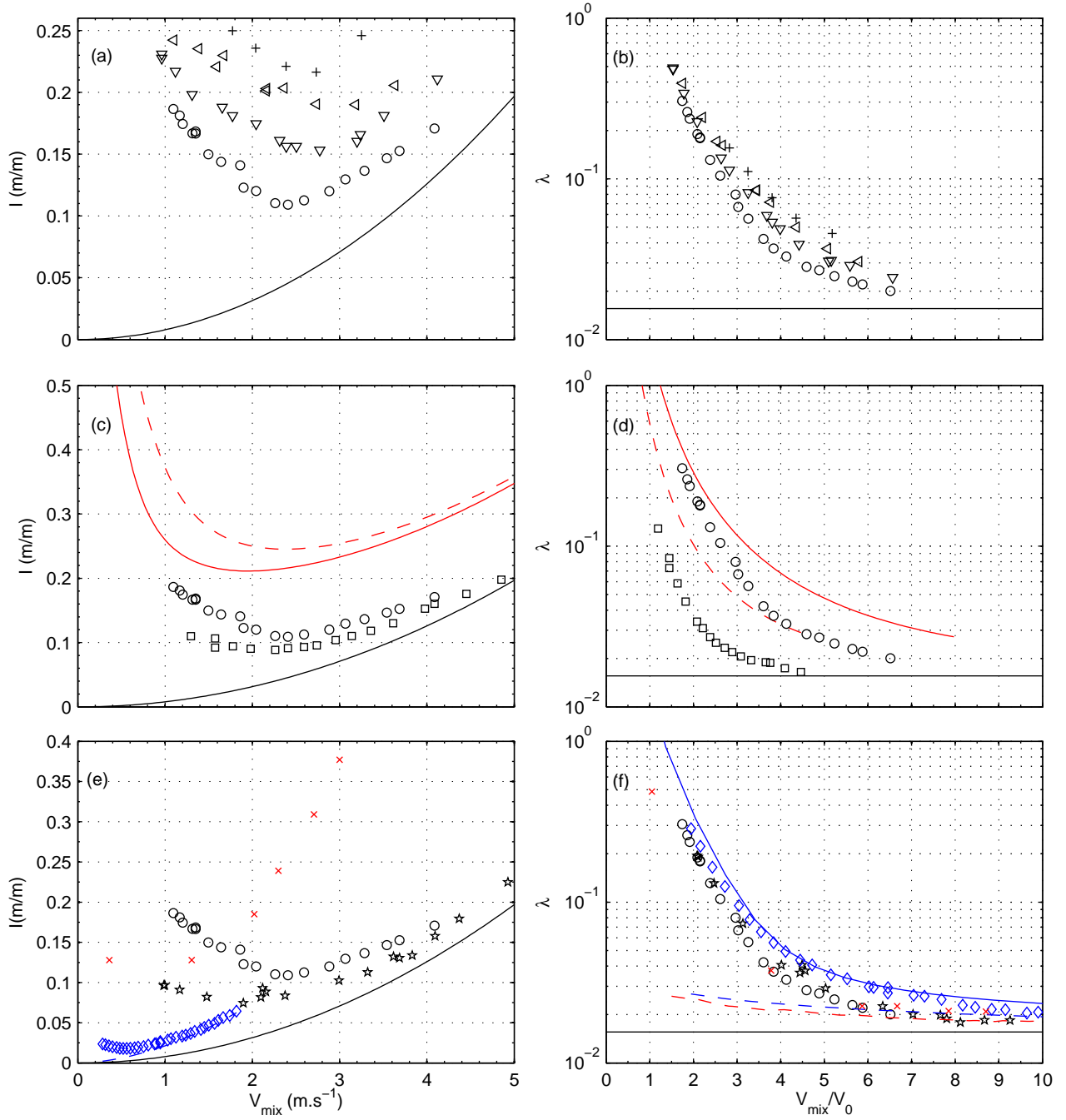


Figure 4: (a,c,e): Hydraulic gradient I_h vs. mixture velocity V_{mix} . (b,d,f): dimensionless plot of λ_h vs. V_{mix}/V_0 . (a-b): Experimental data for Alumina beads of $d_p = 6$ mm at various concentrations: $C = 5\%$ (o), $C = 10\%$ (∇), $C = 15\%$ (\triangleleft) and $C = 20\%$ (+). Solid black line: water flowing alone. (c-d): Experimental data in the horizontal pipe (symbols) and predictions of the model presented in § 3.1 for a similar vertical pipe (red lines), for Alumina beads at $C = 5\%$ and two sizes: $d_p = 6$ mm (o and solid red line) and $d_p = 15$ mm (\square and dashed red line). (e-f): Experimental data for $d_p = 5\&6$ mm and $C = 5\%$ for two specific masses: Alumina (o) and Glass (\star). Blue \diamond : data extracted from Fig. 3 of Ref. [12] (particles of diameter $d_p = 3$ mm, $s = 1.24$ flowing in an horizontal pipe of diameter $D = 50$ mm at $C = 4.2\%$), solid blue line: validation of the two-layer model, dashed blue line: single phase pressure drop. Red \times : data extracted from Fig. 12 of Ref. [4] (gravels of mean diameter $d_p = 2.6$ mm, $s = 2.55$ flowing in an horizontal pipe of diameter $D = 25.4$ mm at $C = 5\%$), red dashed line: single phase pressure drop.

15 mm. The main effects of the size are first that the pressure drop is decreased for the large particles in the horizontal pipe, and secondly that the critical velocity does not seem to be affected by the particle size. This feature is quite surprising to the best of our knowledge has not been reported in previous works that mainly dealt with particles below 4 mm (see § 3.3). It is moreover robust as it is confirmed on the Glass beads (see ★ and ◊ in Fig. 8). The decrease of pressure drop with larger particles and the constancy of the critical velocity are a distinguishing feature between horizontal and vertical flows: owing to the dependence of the slip velocity on the square root of the particle diameter, the hydraulic gradient and the critical velocity are greater for larger solids in vertical flows as a direct result of the model presented in § 3.1.

Finally, the experimental measurements of the pressure drop for two beads of similar size ($d_p \simeq 5$ mm) and two different specific masses, respectively $s = 3.65$ (◊) and $s = 2.5$ (★) are plotted in Fig. 4e-f. Increasing the specific mass leads to both an increase of the pressure drop and of the critical velocity. Experimental data on particles of similar size that are available in the Literature [4, 12] are also plotted in this figure. Please note that they have been collected in pipes of smaller diameters. The shape of the present curves is very similar to previously reported works, particularly when presented in a dimensionless form (λ vs. V_{mix}/V_0 , in Fig. 4f). Please nonetheless notice that the single-phase flow pressure drop coefficients are different (black solid line, blue and red dashed lines in Fig. 4f).

4.2. Mixes of beads and rough solids

This paragraph deals with mixtures of spheres and rough stones of arbitrary shapes in order to check to what extent the results obtained for mono-disperse spheres may represent an actual application.

The figure 5 presents the horizontal hydraulic gradient I_h vs. V_{mix} for different mixtures of Alumina beads of diameter 6 mm and 15 mm. Contrary to what one might think *a priori*, the pressure drop of the mixtures is not a simple linear combination of the pressure drop of each bead size: for a 50% of 6 mm mixture (mixture 1) the pressure drop curve coincides with that of the 15 mm beads. The pressure drop is thus low. This effect is even still present for a proportion of 75% of 6 mm beads in the mixture (mixture 2) but only at low mixture velocities corresponding to $V_{mix} \lesssim V_{crit}$, *i.e.* to regimes with a stationary bed. For higher velocities the pressure drop lies between the other two and is closer to the pressure drop of the 6 mm beads.

The figure 6 presents the comparison of $I_h(V_{mix})$ for three mixtures of beads of same size but different specific mass: Glass beads of 5 mm (★), Alumina beads of 6 mm (◊) and mixture 3 (50% Glass / 50% Alumina, ◊). The pressure drop curve for the mixture lies between the two single-type cases and seems to be well modeled by the mean of the two curves: the solid red line in Fig. 6 is a fit

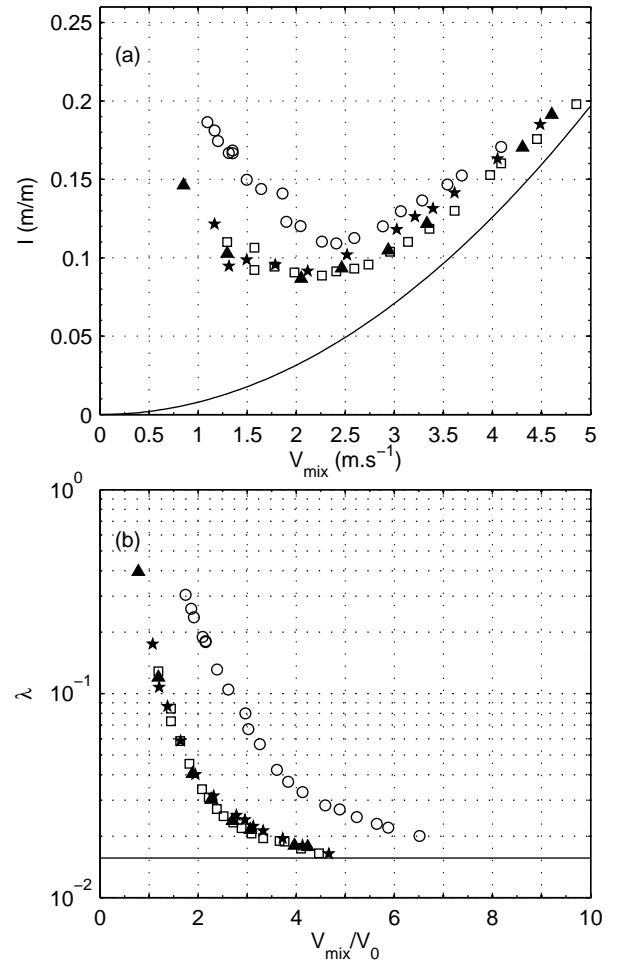


Figure 5: (a): Hydraulic gradient I_h vs. mixture velocity V_{mix} and (b): dimensionless plot of λ_h vs. V_{mix}/V_0 , at $C = 5\%$ vs. for ◊: Alumina 6 mm, ◻: Alumina 15 mm, ◴: mixture 1 and ★: mixture 2.

for the Alumina, the solid blue line a fit for Glass and the black line is the mean of these two curves.

The effects that have been observed for mixtures, particularly in the horizontal part, may be ascribed to segregation phenomena. This is illustrated in Fig. 7 for mixture 2 of Alumina of two sizes (see Tab. 1). A tendency of having two layers of beads, with the small beads being transported at the bottom of the pipe while the large beads are transported on top of this bed, is indeed observed. This phenomenon is reminiscent of the so-called “Brazil nut effect”. Likewise, for the mixture of beads of different specific masses, the heaviest tend to settle at the bottom of the pipe.

All the previous results concern experiments with spherical beads of unique size and specific mass or mixtures of at most two different types of spherical beads. The pressure drop curves for stones are plotted in Fig. 8. Their physical characteristics are given and illustrated in Tab. 1. The specific mass of a sample of fifty solids have been measured with a densimeter. It is constant within $2700 \pm 10 \text{ kg.m}^{-3}$. Their specific mass is thus very close to that of Glass.

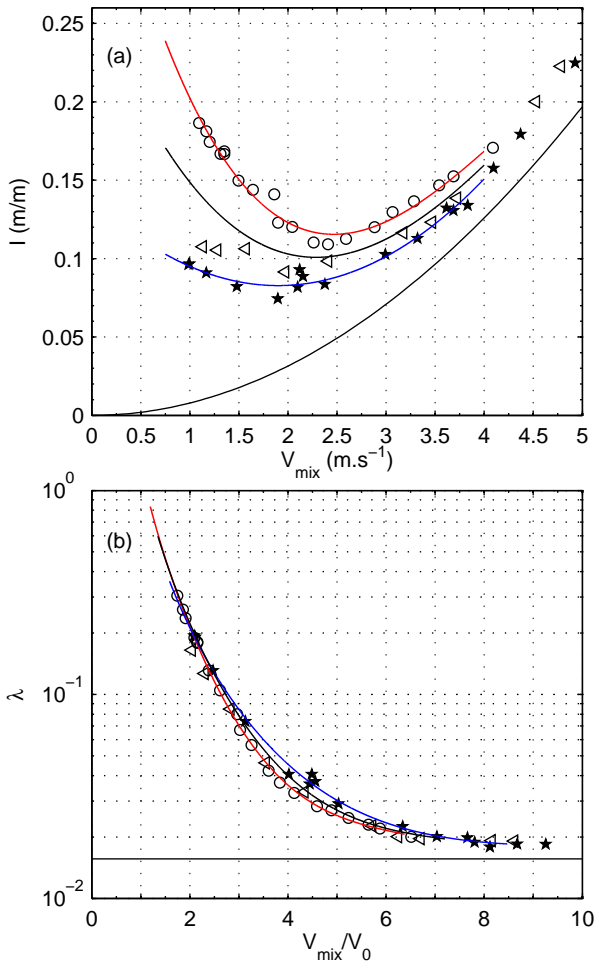


Figure 6: (a): Hydraulic gradient I_h vs. mixture velocity V_{mix} and (b): dimensionless plot of λ_h vs. V_{mix}/V_0 , at $C = 5\%$ for \circ : Alumina 6 mm with a fit corresponding to the red line, \star : Glass 5 mm with a fit corresponding to the blue line and \triangleleft : mixture 3. The black solid line is the mean of the two fits.

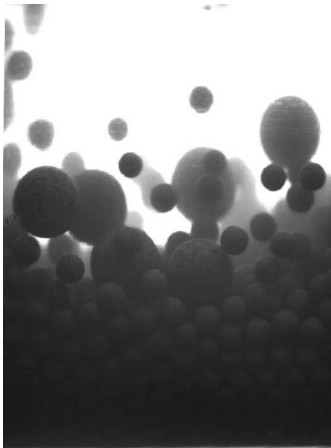


Figure 7: Illustration of segregation for mixture 2. In that case, $C = 5\%$ and $V_{mix} \simeq 1 \text{ m.s}^{-1}$.

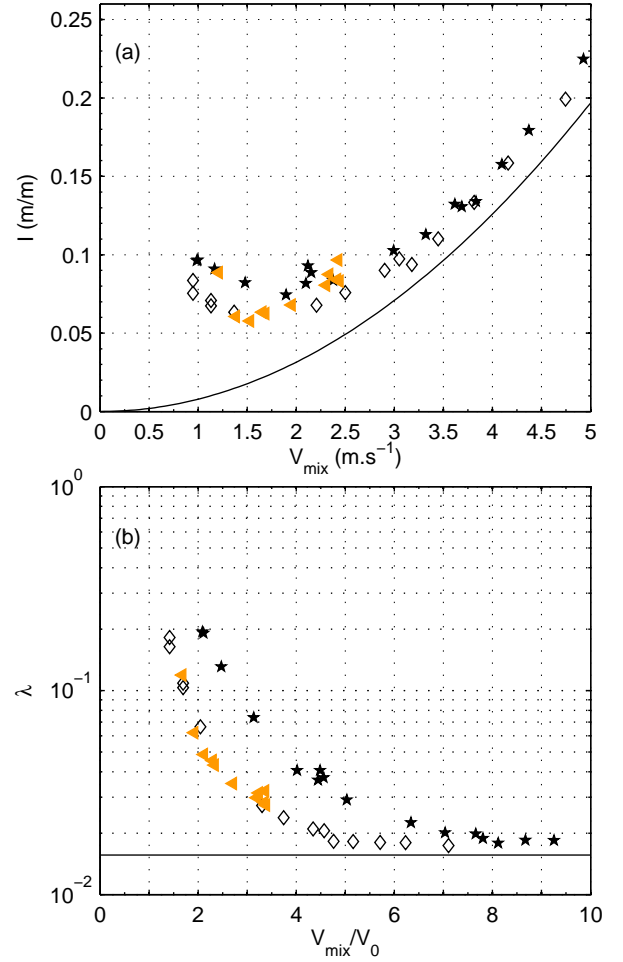


Figure 8: (a): Hydraulic gradient I_h vs. mixture velocity V_{mix} and (b): dimensionless plot of λ_h vs. V_{mix}/V_0 , at $C = 5\%$ for \star : Glass 5 mm, \diamond : Glass 10 mm and orange \triangleleft : stones.

Their size distribution is between 8 and 18 mm, with 50% of the solids having a size lower than 10 mm. The pressure drop I_h for these solids is very close to the case of 10 mm Glass beads. The irregular shape of the solids that leads to different drag coefficients thus does not seem to play an important role with respect to the hydraulic gradient, as already reported by Yoon *et al.* [11]. It may be a second order effect with respect to the size and specific mass effects.

4.3. Analysis and discussion

This section is devoted to the application of various models to the present data and to their discussion.

Semi-empirical correlations. The first quantity that could be checked is the critical velocity V_{crit} that is the mixture velocity for which the minimum hydraulic gradient is observed. The present results suggest that the critical velocity does not strongly depend on the particle size but depends on the specific mass (Fig. 4c-e). The terminal settling velocity furthermore does not seem to be the velocity

scale that drives the pressure drop curve as can be seen in Fig. 4d. The correlations that use the “Froude” number F_D (Eqs. 4 and 5) may be applicable.

The results concerning the prediction of the critical velocity are reported in Tab. 2. The predicted values are in excellent agreement with the experimental values for Glass and Alumina beads of various sizes and at various concentrations (see Figs. 4 for Alumina and 8 for Glass).

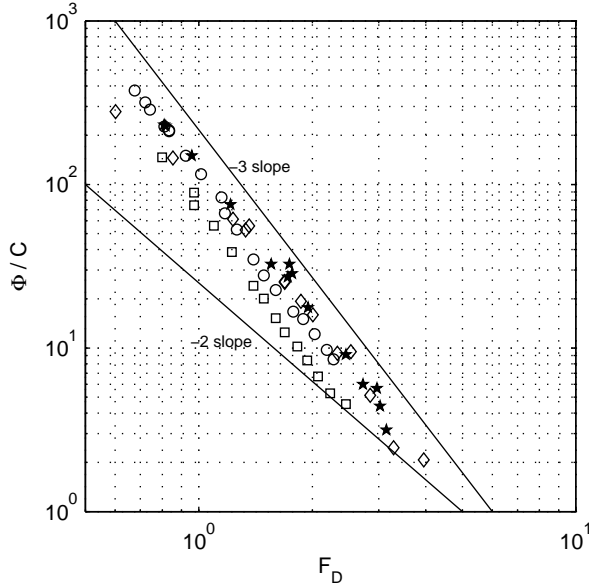


Figure 9: Dimensionless excess of head loss divided by the delivered concentration Φ/C vs. Fr for \star : Glass beads of 5 mm ($C = 5\%$), \diamond : Glass beads of 5 mm ($C = 10\%$), \circ : Alumina beads of 6 mm ($C = 5\%$) and \square : Alumina beads of 15 mm ($C = 5\%$).

The value of the dimensionless excess of head loss Φ , normalized by the delivered concentration C is plotted as a function of the Froude number F_D for various data sets in Fig. 9. All the data follow the same trend. The effect of the concentration is moreover well described by a linear dependence: the data points for Glass beads of 5 mm at a concentration $C = 5\%$ (\star) and at a concentration $C = 10\%$ (\diamond) collapse on a single curve. A -3 power law $\Phi = C K F_D^{-3}$ fits quite well the data in the range $0.7 \lesssim F_D \lesssim 3$. The effect of specific mass that is included in the definition of the Froude number seems to be well taken into account: the data points for Glass beads of 5 mm at a concentration $C = 5\%$, $C = 10\%$ (\star & \diamond) and for Alumina beads of 6 mm at a concentration $C = 5\%$ (\circ) are very close, the values of the constant K being respectively 130 ± 4 and 123 ± 3 . All these considerations are consistent with the empirical correlation (Eq. 5) and the theoretical equation (Eq. 6) with $V_0 \propto \sqrt{s-1}$ for large particles. On the contrary, the -2 power law and the dependency in $(s-1)$ of Φ that are predicted by Eq. 7 does not seem to be consistent with the present measurements. As already noticed, the size of the particles, that is not taken into account in the model, has a strong influence on the hydraulic gradient: the value of the constant for

Glass beads of 10 mm (not represented in Fig. 9) is indeed $K = 87$ and the value for Alumina beads of 15 mm (\square) is $K = 75$. This is predicted neither by Eq. 5 (no dependence in d_p) nor by Eq. 6 (Φ increases as $\sqrt{d_p}$). The model that leads to Eqs. 6 and 7 relies on the hypothesis that $\epsilon_s \propto C$. This is not the case in the present experiments: for Glass beads of $d_p = 5$ mm at $C = 5\%$, the values of the *in situ* concentration estimated with imaging are indeed: $\epsilon_s \simeq 32\%$, 10% and 7% for $V_{mix}/V_{crit,h} = 0.67$, 1.17 and 2.72 . For the Alumina beads of $d_p = 6$ mm and $d_p = 15$ mm, the measured concentrations are $\epsilon_s \simeq 11\%$ and 9% for $V_{mix}/V_{crit,h} = 0.83$ and 1.66 .

Analytical model based on mass and momentum balance. The question that is addressed is to what extent the analytical model of Doron *et al.* [12] may apply to the present physical parameters, with solids of much higher specific mass and even higher relative diameter.

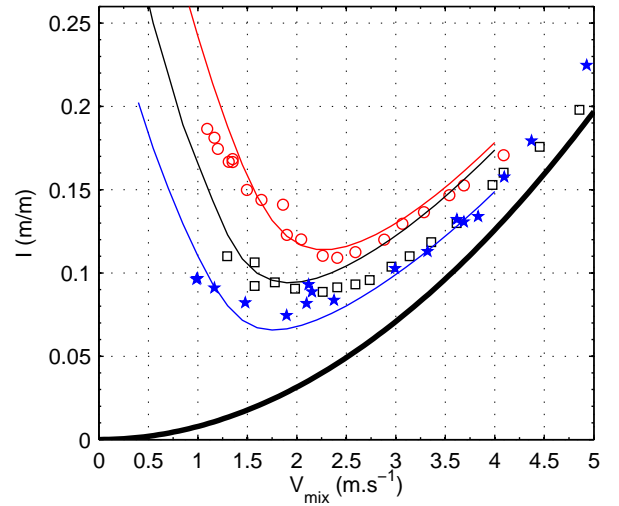


Figure 10: Results of the model (solid lines) compared to the present experimental data (symbols) at $C = 5\%$. Blue line and \star : Glass 5 mm. Red line and \circ : Alumina 6 mm. Black line and \square : Alumina 15 mm. Parameters for the model: $C_b = 0.52$, $\eta = 0.25$, $\tan \varphi = \infty$ (see text).

In the original work of Doron *et al.* [12], the values of the parameters are the following: $\eta = 0.3$, $\tan \varphi = 0.6$ and $C_b = 0.52$. The packing concentration C_b has first been experimentally measured by weighting a tube of same diameter and capacity two liters filled with dry beads and with beads and water. A dozen of measurements have been performed for each type of bead. The concentration is found to be 0.52 ± 0.01 . The determination of the solid friction coefficient for an immersed granular bed is a very difficult problem [20], and the friction coefficient itself can strongly vary with the beads roughness [21]. We thus choose to vary η and to present the results that better match the experimental data in the vicinity of the critical velocity. The angle φ is set to $\varphi = \pi/2$ which corresponds to neglecting the contribution of the normal stress transmitted into the bottom layer by the shear at the interface.

	Glass 5 mm, $C = 5\%$	Alumina 6 mm, $C = 5\%$
Experiment	1.8	2.4
Eq. 4 with $F_l = 1$	1.7	2.3
Eq. 4 with $F_l = 1.05$	1.8	2.4

Table 2: Critical velocity V_{crit} (m.s^{-1}) experimentally measured for Alumina and Glass beads, compared to the values predicted with the correlation 4.

This assumption is validated *a posteriori* by evaluation of its relative contribution.

The results for the hydraulic gradient are presented in Fig. 10. The final value of the friction coefficient is $\eta = 0.25$. The predictions of the model for the two small beads of different specific mass and relative diameter 5% are in relatively good accordance with the experiments. For the largest beads, the trend of the reduction of the hydraulic gradient is reproduced but the model is not satisfactory: it matches the data point only very close to the critical velocity.

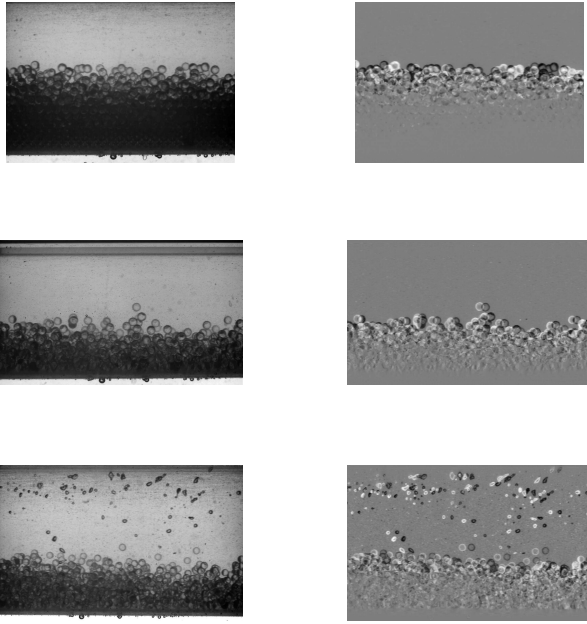


Figure 11: Pictures taken in the horizontal pipe of Glass beads of diameter 5 mm at $C = 5\%$. Left column: rough picture. Right column: difference between two successive images taken at the frame rate f . From top to bottom: $V = 1.0 \text{ m.s}^{-1}$ & $f = 50 \text{ Hz}$, $V = 1.9 \text{ m.s}^{-1}$ & $f = 500 \text{ Hz}$ and $V = 3.9 \text{ m.s}^{-1}$ & $f = 630 \text{ Hz}$.

The model also predicts the height of the lower layer and the velocities of the two layers. Some pictures taken for the Glass beads at $C = 5\%$ are presented in Fig. 11 at different velocities. The right column corresponds to the difference between two successive images and allows to better identify the different flow regimes and the velocity gradient in the bed. At $V = 1.0 \text{ m.s}^{-1}$, one can clearly identify a static bottom layer and a packed layer of 3 to 4 beads that are transported with almost no vertical gradient. The height of the static bed is roughly 42 mm

while the value predicted by the model is 40 mm. At $V = 1.9 \text{ m.s}^{-1}$, the bottom layer is moving slowly with almost no vertical gradient of velocity. The height of the moving packed bed is roughly 25 mm while the value predicted by the model is 22 mm. Above this moving bed, an unpacked layer of suspended beads is observed. And finally, at $V = 3.9 \text{ m.s}^{-1}$, one can still distinguish two layers of solids, with a bottom layer that is moving more rapidly. The height of this static bed is roughly 22 mm while the value predicted by the model is 15 mm. The velocities that are evaluated with the movies give similar accordance.

In conclusion, this two-layer model, originally validated only on small specific mass particles, is promising and predicts well the global and few local features of the two-phase flow for larger specific masses. It seems however limited to not too large particles. One question is the validity of the advection-diffusion equation that is used to determine the concentration in the upper layer for particles that become larger than the turbulent eddies.

5. Conclusion

The hydraulic gradient as a function of mixture velocity has thus been measured for calibrated beads of two specific masses (2500 kg.m^{-3} and 3650 kg.m^{-3}), with large particle to pipe diameter ratios of 5, 10 and up to 15%, as well as for mixtures of calibrated beads and for real stones of similar physical properties. The tests have been conducted in an horizontal pipe. The main results are the following:

- The increase of friction in an horizontal pipe is quite large at velocities of the order of the terminal velocity of a settling particle. This increase of the hydraulic gradient is however lower than that which would be observed because of the hydrostatic contribution in an ascending vertical pipe. On the other hand, the critical velocities that are a key parameter with respect to plugging for the design of a complex industrial application are greater.
- For a given specific mass and delivered concentration, the hydraulic gradient decreases with the increase of the particle size in an horizontal pipe, contrary to what is observed in vertical pipes.
- The empirical correlations that are available in the Literature give satisfactory results but the classical

constants that are recommended clearly do not correspond to the present case of very large particles. Further studies with even larger beads are necessary to better model the variation of these constants with the particle size.

- The mixtures and real stones could be modeled with mono-dispersed beads of equivalent specific mass and size. However, at low velocities, strong segregation mechanisms are present and would make the modelisation harder.
- Analytical models based on more physical arguments are of great interest. They however show their limits for the largest beads.

This last point may be linked to the fact that for large beads the modelisation of the solids using a continuous medium even with the help of granular theory becomes very untrustworthy. The number of particles that can be put in the pipe cross-section is too small to be treated with statistical methods. For instance, numerical methods with Eulerian-Eulerian formulation such as the one presented in Ref. [22] is inappropriate to our case and even give unrepresentable results. Full Lagrangian methods would also be very expensive owing to the nevertheless large number of particles. We are now developing two alternate methods: one is based on Lagrangian dynamics for the solids, the forces being prescribed with the help of a two-layer global model for the fluid. The idea is to detect the position of the compact bed and to compute the concentration in the upper layer, and then to apply some bed-load sediment transport equations [19] for the momentum balance in the bed. The second is to develop a Navier-Stokes code that uses volume penalization [23] that seem very promising to cope with the solids.

Further experiments are scheduled in the S-shape part, and in inclined pipes.

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