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MRI investigation of granular interface rheology
using a new cylinder shear apparatus

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Abstract
The rheology of granular materials near an interface is investigated through proton magnetic resonance imaging. A new cylinder shear apparatus has been inserted in the MRI device, which allows the control of the radial confining pressure exerted by the outer wall on the grains and the measurement of the torque on the inner shearing cylinder. A multi-layer velocimetry sequence has been developed for the simultaneous measurement of velocity profiles in different sample zones, while the measurement of the solid fraction profile is based on static imaging of the sample. This study describes the influence of the roughness of the shearing interface and of the transverse confining walls on the granular interface rheology.

Key words: velocimetry, granular material, annular shear, interface, roughness

1. Introduction
The interaction of a granular material with a solid interface is of interest in various engineering problems, such as industrial conducts [1], geotechnics [2, 3], and in geophysical situations, such as tectonophysics [4] or gravity flows [5].

At the immediate vicinity of the shearing interface, a thin granular layer, where the shear and dilation is localized, plays a significant role in the stress transmission between the solid interface and the bulk granular material. This rheology is influenced by the roughness of the shearing surface [6-7, 10].

In this paper, we focus our attention on the annular (Couette) shear geometry, where the material is confined between two cylinders and sheared by the rotation of the inner rough one (see [3] for a recent review). This geometry has been used to measure the rheological properties of granular materials, both in two dimensions [11-13, 16, 17] and three dimensions [13-15, 20, 22-24, 25, 26, 28]. However, the visualization of the granular interface is usually limited to the upper (free surface) or bottom layers (through a transparent glass window) [13, 17]. Following previous magnetic resonance imaging investigation of granular rheology [30, 31, 32, 33]: flows in rotating drum [34, 35, 36, 37], vertical chute [38], annular shear cell [39, 20, 25], segregation and convection under vibration [40, 41, 42, 43, 44, 45, 46, 47], we have used MRI to measure the granular rheology (velocity and solid fraction profiles) well inside the sample.

Sec. 2 is devoted to the description of a new annular shear cell, specially designed to be inserted in a MRI device. Based on a geotechnical cylinder shear apparatus [19, 48, 49, 7, 29, 50, 51], its originality relies on the control of the radial confining pressure exerted by the outer wall on the grains and on the measurement of the torque on the inner shearing cylinder.

Sec. 3 explains the multi-layer MRI velocimetry. MRI velocity measurements were performed using a spin-warp / phase encoding technique previously adapted from [52] and used for routine liquid rheology in annular Couette cells (see for instance [53, 54, 55, 56, 57]). It was here further modified on purpose of quasi-simultaneous assessment of different regions in the sample.

Sec. 4 describes the measurement of the velocity and solid fraction profiles, as well as the shear stress on the shearing wall, from which we deduce the influence of the roughness of the shearing interface and of the trans-
verse confining walls on the granular interface rheology. Our study is restricted to the quasi-static regime (small shear velocity and/or high confining pressure).

Preliminary and complementary results are presented in [7].

2. Annular shear cell

The annular shear cell is inserted in the MRI device through an external support. The figure shows a schematic view of the cell, which components are made of PMMA.

![Schematic view of the shear cell.](image)

The sample has a hollow cylinder shape. The granular material is confined between two fixed horizontal (bottom and upper) plates (height \( H = 10\, \text{cm} \)), an internal rotating cylinder (radius \( R_i = 3\, \text{cm} \)) and an external latex homemade dip molded membrane (radius \( R_e = 6\, \text{cm} \)) filled with water. A pressure-volume controller (GDS®) applies a radial confining pressure \( P \) to the sample, in the range \( 0 - 15\, \text{kPa} \). An optical fiber sensor (FOP MEMS 1000 kPa - Fiso Technologies®) precisely measures this pressure close to the membrane.

The internal rotating cylinder is guided by two journal bearings set in the external support, while the main cell (membrane, bottom and upper plates) is only connected to the external support through the torque sensor, which measures the whole torsion effort.

An aluminum-alloy torque sensor was specially designed, based on resistive strain gages, in the range \( \pm 10\, \text{Nm} \). This measurement was not possible during MRI experiment, but when displacing the cell bellow the MRI device. This prototype is a first step toward the realization of a torque sensor working inside the MRI device.

Depending on the way upper and bottom plates are mounted, it is possible to measure the whole torsion effort or only the fraction transmitted to the lateral membrane (the difference between those two measurements provides the torque transmitted by the horizontal walls).

The cell is connected to the transmission axis of a rheometer previously designed to be inserted within the MRI facility [58, 56, 59, 60], through a gearbox. This two-stage, timing-belt and pulley system is placed close to the cell and far from the motor. Its reduction factor of 10 provides a rotation range \( 1/600 \leq \Omega \leq 1/6 \, \text{RPS} \).

This configuration allows to place down the cell (out of the MRI tunnel) during the sample preparation and torque measurement and then move it up to the observation position.

The complete cell has a total diameter of \( \approx 19.5\, \text{cm} \) and a total height, without gearbox, of 28 cm, which fits inside the RF coil.

3. Multi-layer MRI velocimetry

MRI experiments were carried out on a Bruker Biospec 24/80 MRI facility operating at 0.5\( T \) (21 MHz proton frequency). The MRI magnet is a vertical superconducting prototype (Magnex Scientific, Oxford), with a 40 cm bore. These characteristics are particularly suited for the study of large and inhomogeneous samples, exhibiting strong internal susceptibility contrasts. The magnet is equipped with a birdcage RF coil (height: 20 cm, inner diameter: 20 cm, hard \( \pi/2 \) pulse duration: 100\( \mu\text{s} \)), and a BGA26 shielded gradient system (Bruker), delivering a 0.05\( \, \text{T/m} \) gradient strength with a rising time of 500\( \mu\text{s} \).

MRI methodology for radial velocimetry inside the cell was that of [52] as further modified by [59] and [60]. It is based on a two-pulse spin echo sequence (Fig. 3a), in which the two pulses are made space-selective in \( z \) and \( y \) direction respectively so as to virtually cut a beam along one cell diameter (Fig. 3b).

A read-out gradient in \( x \) directions permits to get after Fourier transform of the signal a 1D magnetization profile along the beam. An additional pair of gradient pulses in \( y \) direction (in black) induces on the magnetization profile an \( x \) dependent phase shift reading:

\[
\varphi(x) = \gamma G\delta \Delta v_y(x),
\]

(1)
where $\gamma$ is the gyromagnetic ratio of proton, $G$ the strength of these 'velocity' pulses, $\delta$ and $\Delta$ are timing components of the sequence as described in Fig. 3, and $v_r(x)$ is the $y$ velocity component along the selected diameter. In order to get a velocity profile, one performs two MRI measurements with respectively positive and negative velocity gradients, and then compares in each pixel the phase of the two magnetization profiles. When the thickness of the beam in $y$ direction is small enough as compared with the cell diameter, such measurement may be regarded as a direct measurement of the orthonal-normal radial velocity component $v_{\theta}$ versus the radial coordinate $r$ (Fig. 3).

![Figure 2: (a) MRI sequence used for velocity measurements, and its coherence pathway. Velocity encoding gradients are shown in black. (b) Cell scheme and virtual beam cut.](image)

In this sequence, only the phase of NMR signal bears relevant information. It is then safe to use recycling times $TR$ equal or even shorter than $T_1$ values. Indeed, an incomplete spin-lattice relaxation between two successive sequences may only affect the signal to noise ratio, but is not prone to introduce any bias, provided that the magnetization be correctly cleaned from past solicitations before each sequence. Such cleaning was here performed with a saturation module at the beginning of the $TR$ delay.

Thanks to saturation, $TR$ may even be chosen so as to optimize the signal to noise ratio of the experiment. Let’s imagine that, because of experimental constraints, one has a fixed delay $T_{exp}$ to perform a velocity measurement. If some $TR$ delay is used, then it will be possible to repeat the sequence $N = T_{exp}/TR$ times. Granted that the available magnetization at the beginning of each sequence scales as $1 - \exp(-TR/T_1)$, and that the improvement of signal to noise ratio when repeating sequences scales as $\sqrt{N}$, the signal to noise ratio $SNR$ of the whole measurement process scales as:

$$SNR(TR) \propto \left(1 - \exp\left(\frac{TR}{T_1}\right)\right) \sqrt{\frac{T_{exp}}{TR}}$$  \hspace{1cm} (2)

This quantity is maximum for:

$$TR_{opt} = 1.26T_1,$$  \hspace{1cm} (3)

and fairly stays above 90% of this maximum in the range $0.57T_1 \leq TR \leq 2.59T_1$. Saturation then turns out a useful tool for signal optimization.

For the assessment of the velocity profiles at different height in $z$ direction, sequences were repeated while systematically shifting the vertical position of the beam. In order to study unsteady granular flows, it was important to make these multi-layer measurements as simultaneous as possible.

First of all, looking back at the NMR sequence, the measurement of one beam at a given position clearly affects all the magnetization of a larger region in the cell, composed of the sum of the two slices selected by each RF pulse. This, together with the use of the saturation module, prevents considering measurements at different $z$ positions as independent: it is not possible to run the sequence at one height immediately after running the sequence at another height, even if the two associated beams are well separated. Real simultaneous measurements of velocity in multiple layers were then not possible.

We proceeded instead as follows (Fig 3). The whole measurement consisted in two nested loops. In the first loop the vertical location of the beam was shifted after each single sequence, so as to go through all necessary beam positions. The procedure was then just repeated with inverted velocity gradient strength. This first loop was inserted in a second one consisting in standard phase cycling and signal accumulation. One could get this way velocity profiles at different height measured over exactly the same period of time.

In the present work, the thickness $t$ of the beam in $y$ direction was 10 mm, and the recycling time $TR$ was...
Figure 3: Organization of MRI velocimetry sequences for the quasi-simultaneous assessment of n layers in the cell.

4. Experimental results

Three roughness levels of the rotating cylinder are tested: the first one corresponds to a vertically corrugated surface (triangular shape with thickness equal to depth of 1 mm), the second one to a glued seeds surface and the third one to sandpaper (medium grain - 80). The horizontal walls (upper and bottom) are originally smooth. To modify their roughness we have sticked sandpaper and glass beads with the same diameter as the seeds (for practical reasons the beads are glued to sandpaper and not directly over the plate surface).

Mustard seeds are used as model granular material (mean diameter \(d = 1.5 \text{ mm}\), quite monodisperse, mass density of 1200 kg/m\(^3\)). The relaxation times are \(T_1 \approx 100 \text{ ms}\) and \(T_2 \approx 40 \text{ ms}\).

The granular material is poured inside the cell with a funnel from the top and then levelled off (see Fig. 4). After closing the cell, the sample is confined by the application of the radial pressure through the membrane. Then, the material is pre-sheared (at least 15 rotations of the cylinder, corresponding to \(\approx 2800 \text{ mm}\) of tangential displacement of the wall) to drive it in a steady-state. Then data acquisition starts.

Figure 4: (Color on line) Sample preparation.

Table 1 summarizes the parametric study (wall roughness, confining pressure \(P\) and velocity at the wall \(V_0 = 2\pi R \Omega\)).

<table>
<thead>
<tr>
<th>cylinder</th>
<th>plates</th>
<th>pressure(^1) (kPa)</th>
<th>velocity(^2) (mm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>corrugated</td>
<td>smooth</td>
<td>3.5 – 13.5</td>
<td>0.314—3.14</td>
</tr>
<tr>
<td>glued seeds</td>
<td>smooth</td>
<td>3.5 – 13.5</td>
<td>0.314—3.14</td>
</tr>
<tr>
<td></td>
<td>sandpaper</td>
<td>8.5</td>
<td>3.14</td>
</tr>
<tr>
<td></td>
<td>glass beads</td>
<td>8.5</td>
<td>3.14</td>
</tr>
<tr>
<td>sandpaper</td>
<td>smooth</td>
<td>3.5 – 13.5</td>
<td>0.314—3.14</td>
</tr>
</tbody>
</table>

\(^1\) pressure (kPa) \(^2\) velocity (mm/s)

Table 1: List of velocimetry experiments.

Internal stresses associated to gravity (of the order of 0.7 kPa) can be considered to be dominated by the applied confining pressures.

The maximum value of the inertial number \([3]\), used to qualify the granular flow regime, can be estimated around \(3 \cdot 10^{-4}\), indicating that the granular material is in the quasi-static regime.

We now discuss the radial profiles of velocity and solid fraction. They are presented as a function of \(r/R_i\), normalized by the diameter \(d\) of the grains.
4.1. Velocity profiles

We have first measured the profiles of the orthoradial velocity \(v_\theta(r)\), normalized by \(V_\theta\), far from the horizontal walls, comparing three layers 15 mm thick separated by 10 mm (Fig. 5).

Figure 5: Scheme of layers in the central zone.

Fig. 5 shows that \(v_\theta(R_i) < V_\theta\), which signifies that there is significant sliding at the wall. Fig. 6 also shows that there is no influence of the roughness of the horizontal walls in the central zone. An approximately exponential decay of the orthoradial velocity profiles \(v_\theta(r)\) is measured, with a maximum value at the shearing wall, consistently with previous studies [18, 19, 20, 14, 29, 13, 15, 25]. Qualitatively, this shear localization is explained by the strong decrease of the shear stress away from the inner wall, while the normal stress remains approximately constant. As expected in the quasi-static regime, no influence of the velocity at the wall \(V_\theta\) nor of the confining pressure \(P\) has been observed.

4.1.1. Influence of the roughness of the horizontal walls

In a 2D system, the shear stress is entirely transmitted from the inner to the outer wall by the granular material [2, 3]. In a 3D system, the granular material interacts with the horizontal walls, to which part of the torque is transmitted. This changes the stress distribution within the sample, and may consequently affect the 3D velocity profile. This influence of frictional lateral walls has been evidenced in granular surface flows between vertical walls [61], especially when they are rough [62]. We now wonder if such effect is observed close to the horizontal walls.

The analysis of the influence of the horizontal walls was based on a comparison of the profiles of 5 layers 1.5 mm thick separated by a distance of 1.75 mm (Fig. 5). The gap of 0.45 mm between the horizontal wall and the first layer ensured that only moving particles were considered in the measure. In the rough case, a similar gap was taken from the glued glass beads surface giving a total thickness of 2.5 mm.

The horizontal roughness made of glass beads is not detectable by MRI, which provided a correct evaluation of the velocity of the grains in direct contact with this surface. The motion of the grains could be clearly distinguished from the motion of the cylinder, given the strong velocity gradient.

Fig. 8 shows a comparison between the profiles in the central zone and near the horizontal wall. In the case of a smooth horizontal wall, there is no significant influence. This result has practical applications, validating the measure of displacements of granular materials through transparent glass walls [19, 7, 29]. Conversely, in the case of a rough horizontal wall, a signif-
ificant decrease is observed very close to the horizontal wall, particularly in the first layer. This indicates the transmission of shear stress between the granular material and the horizontal walls. However, the perturbation remains very localized, in contrast with granular surface flows [32]. The fluctuations of the velocity profiles in the first layer close to the horizontal wall is explained by the small thickness of those layers, which strongly decreases the signal to noise ratio.

4.1.2. Influence of the cylinder roughness

Fig. 8 shows that the cylinder roughness has a significant influence on the velocity $v_y$, causing offsets of the whole velocity profiles. Sliding, defined as the ratio between the maximum value of the particle velocity and the wall velocity $V_0$ [29, 50, 63], decreases from sandpaper cylinder to the corrugated one, that is to say as the roughness increases.

4.2. Friction coefficient

The average shear stress at the inner cylinder is simply deduced from the measured torque $C$ as $S = C/(2\pi R^2 H)$. Consistently with previous studies [29, 35, 50], a linear relation between the shear stress $S$ and the confining pressure $P$ is observed (Fig. 3), from which an effective friction coefficient $S/P$ is deduced (considering that the normal stress at the cylinder wall is approximately equal to $P$) for each of the three cylinders: 0.35 for sandpaper, 0.38 for glued seeds and 0.4 for corrugation. This increase seems consistent with the decrease of the particle sliding shown in Fig. 8. The roughness of the horizontal walls does not affect this value, which is consistent with the very localized influence of this roughness.

4.3. Solid fraction profile

In order to measure the influence of the shear on the radial distribution of grains, 3D spin density pictures (20 layers 5 mm thick in the $z$ direction, with a space resolution of 0.2 mm in both $x$ and $y$ directions) were performed at rest before and after shearing (at least 200 rotations, making $\approx 38$ m of tangential displacement).
5. Conclusions

As a way to study the rheology of granular materials near an interface, we have developed a cylinder shear apparatus inserted in the MRI device, which allows the control of the radial confining pressure exerted by the outer wall on the grains and the measurement of the torque on the inner shearing cylinder. We have also developed a multi-layer velocimetry sequence for the simultaneous measurement of velocity profiles in different sample zones. This study shows that the roughness of the shearing interface significantly affects the granular interface rheology (sliding velocity and friction coefficient), while the influence of the roughness of the transverse confining walls remains localized in the very first layers. Moreover, the measurement of the solid fraction profiles, based on static imaging of the sample, shows that, when starting to shear the material, its volumetric variations remain localized close to the shearing interface. These data are useful to test models attempting to describe granular interface rheology for annular shear. Using this device together with the velocimetry sequence, work is under progress to study the transient behavior of the granular material, under...
monotonic or cyclic shear, and the influence of liquid saturation.

References


