

Stretching and folding sustain microscale chemical gradients in porous media

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Fluid flow in porous media drives the transport, mixing, and reaction of molecules, particles, and microorganisms across a wide spectrum of natural and industrial processes. Current macroscopic models that average pore-scale fluctuations into an effective dispersion coefficient have shown significant limitations in the prediction of many important chemical and biological processes. Yet, it is unclear how three-dimensional flow in porous structures govern the microscale chemical gradients controlling these processes. Here, we obtain high-resolution experimental images of microscale mixing patterns in three-dimensional porous media and uncover an unexpected and general mixing mechanism that strongly enhances concentration gradients at pore-scale. Our experiments reveal that systematic stretching and folding of fluid elements are produced in the pore space by grain contacts, through a mechanism that leads to efficient microscale chaotic mixing. These insights form the basis for a general kinematic model linking chaotic-mixing rates in the fluid phase to the generic structural properties of granular matter. The model successfully predicts the resulting enhancement of pore-scale chemical gradients, which appear to be orders of magnitude larger than predicted by dispersive approaches. These findings offer perspectives for predicting and controlling the vast diversity of reactive transport processes in natural and synthetic porous materials, beyond the current dispersion paradigm.

porous media | reactive transport | chaotic mixing | chemical gradients

Fluid mixing in porous media plays a key role in a range of natural and industrial systems (1–3). In these confined environments, mixing enables or limits reactions controlling the degradation of contaminants in the subsurface; the cycles of biogeochemical elements such as nitrogen, iron, and carbon; and the sequestration of CO_2 in deep reservoirs (4–10). Mixing also shapes the nutrient landscapes and chemical gradients seen by bacteria evolving in soils or medical systems (11, 12) and facilitates chemical processes in drug delivery, packed bed reactors, flow batteries, or catalysts (13-15). Increasing evidence of sustained chemical gradients and incomplete mixing below the pore-scale, along with associated impacts upon chemical reactions (3, 16–18), have questioned the relevance of macroscopic dispersion coefficients to capture these processes (1). Yet, it is currently unknown how three-dimensional flow topologies in porous structures control microscale mixing rates and concentration gradients.

Recent theories (19, 20) have suggested that laminar flow through three-dimensional porous media may possess the basic ingredients for chaotic advection (e.g., the exponential deformation of fluid elements), which would represent a possible mechanism for the enhancement of microscale chemical gradients and the persistence of incomplete mixing at the pore-scale. These chaotic dynamics may have particularly important consequences for microbial processes, a broad range of which are hosted in porous environments (21). Biological processes in turbulent flows have been shown to be deeply altered by chaotic advection, which promotes coexistence of competitive microbial species (22) and affects the chemotactic responses of microor-

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ganisms (23). However, whether such chaotic dynamics can spontaneously develop in laminar flows through porous media remains on open question.

A key experimental barrier to the direct imaging of solute advection in three-dimensional porous materials is their predominantly opaque nature. While X-ray microtomography technologies have progressed significantly (24), they still cannot resolve the fine structures produced below pore-scale. In contrast, use of visible-spectrum refractive-index matching between the solid and the fluid phases represents a viable alternative to observe solute mixing, as obtained with hydrogel beads in water (25). However, as molecular diffusion eventually masks the deformation of dved fluid elements, a direct measurement of fluid deformation in random porous media is an outstanding challenge. Here, we overcome these limitations by performing high-resolution laser imaging of the evolution of a low-diffusivity fluorescent dye plume through a column of optically transparent borosilicate spheres via high-precision refractive-index matching (Fig. 1). This technique allows reconstruction of the three-dimensional dye plume at unprecedented resolution, thus providing direct experimental observation of pore-scale fluid deformation and mixing in porous media. These data reveal the hitherto unknown role of grain contacts in controlling folding and stretching of fluid elements, a mechanism that generates strong chaotic advection and significantly enhances chemical gradients at the microscale. Since grain contacts are inherent to all

Significance

Porous media flows are central to environmental, industrial, and biological systems, where they transport molecules, particles, and microorganisms and trigger important biogeochemical reactions. There is increasing evidence that many of these processes are highly sensitive to chemical gradients below the pore-scale. However, it is unknown how porous architectures control microscale concentration heterogeneities. Here, we provide high-resolution experimental images that fully resolve three-dimensional pore-scale mixing dynamics. We show that grain contacts control the folding of fluid elements in the pore space, which, in addition to fluid stretching at stagnation points, leads to the exponential enhancement of microscale concentration gradients. We use these insights to derive a kinematic model linking mixing rates to pore geometry, opening perspectives for reactive transport modeling.

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Fig. 1. (*A*) Experimental dye plume with backbone filament (red lines) at selected stages of the folding mechanism (run III; Movie S4). Beads have been shrunk by 40%, and white lines depict the contact line joining bead centers; surrounding beads are not shown. (*B*) Cross-sections detailing the typical folding stages (run I; Movie S3). Upon passing the contact point between beads 1 and 2 (*B.0*), the filament is stretched (diverging arrows) along the contact plane (dashed line) and compressed in the perpendicular direction (converging arrows). After contact (*B.1*), the directions of stretching and compression are inverted, and a cusp forms, creating a fold (*B.2*) that is advected over the subsequent contact between beads 3 and 4 (*B.3*). Other experimental runs and numerical simulations are reported in *SI Appendix*, Figs. S1 and S2 and Movie S5.

granular porous materials, we deduce that chaotic mixing is ubiquitous in flow through all such materials, potentially impacting a large range of fluid-borne phenomena in natural and engineered systems.

Three-Dimensional Imaging of Mixing Patterns in Porous Media. We observed three-dimensional fluid deformation and solute mixing in laminar flows through monodispersed random bead packs of diameters d = 7, 10, and 20 mm, optically matched into a glycerol-water fluid mixture (see Fig. 5). A fluorescent dye is continuously injected upstream of the transparent column as a thin tube of radius $L_0 \ll d$ (Fig. 2A) and advected downstream by the porous flow at the mean longitudinal advection velocity *u*. Cross-stream concentration patterns of the dye plume are imaged in the pore-space via a translational scan using a laser sheet and a camera. The dye cross-section rapidly evolves into a highly elongated (Fig. 2B) and striated filamentous structure (Fig. 2C) due to transverse stretching and folding of fluid elements in pores (Movies S1-S3). The combination of a highly viscous fluid mixture and a high-molecular-weight dye results in laminar flows of low diffusivity, characterized by Reynolds numbers on the order of Re = $7 \cdot 10^{-3}$ and Péclet numbers on the order of $Pe = 10^4$ (*SI Appendix*, Table S1). The deformation of the dye plume (Fig. 1A) thus closely shadows that of the advected fluid, facilitating direct visualization of pore-scale fluid deformation. We use spline fitting on the images to reconstruct the backbone of the cross-sectional dye footprint, called a filament, and estimate its total length L(x) for 9 to 14 bead diameters downstream from the injection point (Fig. 2 and Movie S3).

The mean total filament elongation L/L_0 , averaged over the 18 statistically equivalent packings (SI Appendix, Table S1),

exhibits clear exponential growth with normalized longitudinal distance x/d (Fig. 3). The dimensionless exponent $\mu \equiv \ln(L/L_0)/(x/d) = 0.29 \pm 0.01$ is independent of both bead diameter and flow rate and is known as the *topological entropy* of the flow (26). Via the central limit theorem, μ is related to the mean λ and variance σ_{λ}^2 of the dimensionless stretching rate (*Materials and Methods*)

$$\mu = \lambda + \sigma_{\lambda}^2 / 2.$$
 [1]

The dimensionless parameter λ is also known as the Lyapunov exponent (26), which can be converted into a mean stretching rate per unit time as $\lambda u/d$. In addition to stretching, the filament also undergoes highly localized folding events that result in closely foliated striations (Fig. 2C). These fluid deformations are the hallmarks of chaotic mixing, thus permitting exponential elongation of material elements in a finite-sized domain.

The Role of Grain Contacts in Folding. Folding of dye filaments is consistently initiated downstream of contact points between two beads (Fig. 1*B* and *SI Appendix*, Fig. S1). The cusp-shaped geometry near grain contacts means that when crossing a contact point (Fig. 1, *B.0*), fluid elements are first compressed in the direction joining the two bead centers and stretched in the perpendicular direction. Downstream of the contact point, the direction of compression and stretching are exchanged and a cusp forms locally in the dye filament (Fig. 1, *B.1*). This cusp is stretched in the following pore space, leading to a folded filament made of two straight segments (Fig. 1, *B.2*). This stretching and folding process is repeated sequentially as the folded filament

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Fig. 2. Cross-stream sections of the fluorescent dye plume at increasing distances from the injection: x/d = 0 (*A*), 3.2 (*B*), and 9.3 (C) (run II; Movies S1 and S2). Colors represent local dye concentrations. Beads appear in deep blue. A spline curve (dotted black lines) is used to fit the filament backbone on each cross-section image, from which the total filament length *L* is obtained.

encounters other contact points (Fig. 1, *B.3*), leading to thin solute dye foliations that are the hallmarks of chaotic advection (26) (Fig. 2*C*).

Recent studies (19, 20) identified the role of separation and reattachment points on open grain boundaries (saddle points) in generating exponential stretching of fluid elements. Here, we uncovered the distinct role of contact points between grains in generating systematic folding of fluid elements. Simulations of laminar flow in periodic bead packings (SI Appendix, section A) show that attracting and repelling stream surfaces (unstable and stable manifolds) produced by these saddles indeed control stretching of material lines in the pore space. We found that these manifolds intersect orthogonally at grain contacts (SI Appendix, Fig. S2 and Movie S5), where both the local flow velocity and the stretching rate vanish and manifold stabilities are exchanged, so that repelling stream surfaces become attracting and vice versa. Hence, over a contact point, the local flow structure imparts finite curvature to fluid elements, which results in the sharp folds observed experimentally (Fig. 1). The repetition of this basic stretching and folding sequence over successive contact points offers a simple geometric framework to relate stretching rates to granular structure.

Linking Stretching Statistics to the Porous Structure. Sharp folding of dye filament in between contact points produces a number n_c of localized cusps of strong curvature, separated by straight segments of moderate curvature (see Fig. 6). We define the mean segment length as $L_c = L/n_c$ and the average area swept out by a segment between two successive contact points as $S_c = S/n_c$, where S is the total area swept by the filament. We find that L_c and S_c both converge to stationary values (Fig. 3, *Inset*). Hence, the average advection distance between two successive contact points is statistically constant and equal to $X_c = S_c/L_c \approx 3.45 d$. Over this distance, a single elementary segment folds, giving rise to two new straight elementary segments. Thus, over the same distance, its length $\ell(X)$ must double to maintain $\langle \ell \rangle = L_c$ constant, so that $\ell(X) = 2^{(X/X_c)}$. Hence, the average dimensionless stretching rate λ of elementary fluid segments can be estimated as

$$\lambda \equiv \frac{\mathrm{d}(\log \ell)}{\mathrm{d}(X/d)} = \frac{\log 2}{X_c/d} \approx 0.21.$$
 [2]

This rate is larger than found in synthetic porous media, such as random pore networks ($\lambda \approx 0.12$) (19) and body-centered cubic assemblies of spheres ($\lambda \approx 0.128$) (27), reflecting the remarkably efficient stretching and folding process occurring in random granular media. The variance of the stretching rate can be estimated from Eq. 1 as $\sigma_{\lambda}^2 \approx 0.16$, a value comparable to the mean stretching rates λ , as it is typically the case in space-filling chaotic flows (28).

As shown by Eq. 2, the strength of chaotic advection is entirely governed by the spatial frequency X_c^{-1} with which segments encounter grain contacts. We show in *Materials and Methods* that in isotropic packings, $X_c \approx 8 \log 2\phi z_c d_p/3$, with z_c the coordination number (the mean number of contacts per bead), ϕ the solid volume fraction, and d_p the mean pore diameter. This yields a simple geometric estimate of the dimensionless Lyapunov exponent

$$\lambda \approx \frac{3}{8} \frac{\phi z_c d_p}{d}.$$
 [3]

Eq. 3 is also applicable to nonisotropic packings with a prefactor that quantifies the distribution of orientations of the contact lines joining bead centers with respect to the mean flow direction (*Materials and Methods*). Insertion of the experimental values $z_c = 6$, $\phi = 0.5$, and $d_p/d = 0.24$ in Eq. 3 yields $\lambda \approx 0.27$, which is in reasonable agreement with the experimental estimate of 0.21, given the slight anisotropic nature of our experimental packings (*Materials and Methods*). Eq. 3 provides a quantitative link between microscopic fluid stretching rates and porous media structural properties.

Discussion

Stretching and Folding Sustain Microscale Chemical Gradients. Repeated sequences of stretching and folding leads to exponential compression of fluid elements that can sustain concentration gradients at the pore-scale (Fig. 2). These concentration gradients are locally controlled by the balance between diffusive spreading rate (D_m/s^2) , with D_m the molecular diffusivity,



Fig. 3. Total filament elongation L/L_0 with respect to the normalized pore advection distance x/d from the dye injection point. (*Inset*) Convergence of the mean segment length $L_c = L/n_c$ and area $S_c = S/n_c$ toward constant values, with n_c , the number of cusps.

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and the mean compression rate $(\lambda u/d)$, which is equal to the stretching rate in steady incompressible flows. These two rates equilibrate at the Batchelor scale

$$s_B = \sqrt{D_{\rm m} d / (\lambda u)},$$
 [4]

which is the characteristic length scale of solute-concentration fluctuations. Pore-scale mixing is thus characterized by the dimensionless length scale $s_B^* = s_B/d$, which is related to the macroscopic Péclet number $\text{Pe} = ud/D_{\text{m}}$ as,

$$s_B^* = (\lambda P e)^{-1/2}.$$
 [5]

For $s_B^* > 1$, the length scale of solute-concentration fluctuations are larger than the grain diameter, and therefore concentration fields are expected to be well mixed at the pore-scale. Our estimation of the Lyapunov exponent λ implies that the transition to incomplete pore-scale mixing occurs at Péclet numbers above 5, values commonly encountered in many natural and industrial contexts (29). This suggests that a broad range of biogeochemical processes are possibly affected by incomplete pore-scale mixing and chaotic advection. From results established on chemical and microbial processes in turbulent and chaotic flows at comparable mixing rates (22, 23), it is possible to anticipate a range of possible effects of chaotic mixing in porous media, including altered effective kinetics and microbial growth dynamics, increased biodiversity, and enhanced benefit of sensing. Because they can only develop in three-dimensional topologies under steady conditions (26), these chaotic dynamics are generally absent in quasi two-dimensional microfluidic experiments used to investigate the microscale interactions between flow, concentration gradients, chemical reactions, and microbial processes (3, 11, 30). Novel experiments in three-dimensional porous media and new modeling frameworks are thus needed to explore these dynamics.

Microscale Mixing Model. The experiments in this study have used high Péclet numbers to uncover the rate and kinematics of fluid deformation in porous media. These results may be extended to the prediction of macroscopic mixing rates and concentration statistics at arbitrary Péclet numbers via lamellar mixing models that couple stretching and diffusion (31–33). In SI Appendix, section C, we derive such a mixing model and compare its predictions in terms of dye-concentration statistics to the experimental data. The lamellar model successfully captures the measured exponential decay of the mean maximum solute concentration of dye filaments with longitudinal distance $c_{\text{max}} \sim \exp(-(\lambda + \sigma^2/2)x/d)$ (Fig. 4), as well as the growth of concentration fluctuations caused by variability of the Lagrangian stretching history. These predictions provide an independent validation of the estimated mean and variance of the stretching rate, $\lambda \approx 0.21$ and $\sigma_{\lambda}^2 \approx 0.16$. In contrast, conventional mixing models based upon macroscopic dispersion coefficients (1) ignore incomplete mixing at the porescale and predict an algebraic decay of concentrations $c_{\rm max} \sim$ $(x/d)^{-1/2}$ (SI Appendix, section B). From the normalized Batchelor scale (Eq. 5), pore-scale concentration fluctuations predicted by the lamellar model will persist for all Péclet numbers larger than 5. In this range, macroscopic dispersion models fails to resolve pore-scale concentration gradients, leading to incorrect predictions of a broad range of reactive transport dynamics (3, 16-18). Coupling lamellar mixing models with reactive processes is therefore a promising avenue to capture the effect of pore-scale incomplete mixing on biogeochemical dynamics.

Porous Materials as Mixers. From Eq. 3, the mixing efficiency of steady laminar flows through random bead packs (defined as

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Fig. 4. Experimental average (triangles) and SD (circles) of the local maximum concentration c_{max} along x. Values are normalized by the initial maximum concentration c_0 . Classical model predictions are shown as black dashed and dotted lines. Lamellar mixing model predictions for the average and SD of c_{max} are shown respectively as the green dotted line and the purple continuous line (see *SI Appendix*, sections B and C for derivations).

the ratio of the average stretching rate to the average strain rate) is found to be 3% (SI Appendix, section D). This value is comparable to the performance of industrial mixers (26) and an order of magnitude larger than that of microfluidic chaotic mixers (34), thus opening opportunities for exploiting the mixing properties of porous materials. Chaotic advection is known to both increase dispersion transverse to the mean flow direction and retard longitudinal dispersion (35). It also alters the transport of finite-sized particles such as colloids and microorganisms (36) and may thus control their clustering in the pore space and deposition on grain boundaries. In relating stretching rates to the porous microstructure, Eq. 3 offers a possible pathway to the design of engineered porous materials with optimum mixing characteristics. This concept may find important applications in the design of heat exchangers; packed bed filters and reactors, where transverse dispersion and mixing act to enhance process efficiency; and for continuous flow chemistry (15), such as pressure-driven chromatography, where product selectivity and yield strongly depend upon the minimization of longitudinal dispersion. These applications would first require a validation Eq. 3 over a large range of packing geometries.

Conclusions

Using high-resolution experimental imaging of microscale mixing in three-dimensional granular media, we have demonstrated the existence of efficient stretching and folding of fluid elements at the pore-scale. We use these insights to develop a stochastic model for the prediction of the Lyapunov exponent from the geometric properties of the grain pack and validate this model against experimental observations. The formalization of these observations into a chaotic-mixing model, coupling stretching and diffusion, demonstrates that incomplete mixing persists at pore-scale for Péclet numbers above 5. This model captures the the evolution of microscale chemical gradients, opening perspectives for understanding, predicting, and controlling a large spectrum of physical, chemical, and biological processes, in natural and engineered porous systems.

The discovery of systematic and efficient chaotic mixing in single phase laminar flows through random bead packs—the archetype of porous media—calls for deeper investigation of this

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Fig. 5. Experimental setup for three-dimensional imaging of the mixing of a fluorescent solute dye in optically matched porous bead packs. The dye is continuously injected through a thin needle upstream of the granular column, while maintaining a steady background laminar flow. After stabilization of the solute dye plume, transverse cross-sections of concentrations are obtained by displacing a laser sheet (wavelength, 488 nm; beam waist 50 μ m; Oxxius) in the x direction while recording the emitted fluorescence with a camera (16 bits; Hamamatsu ORCA-Flash4.0) mounted with a bandpass filter (Midopt BN532) and oriented at an angle of 45° from the laser plane. The perspective view is then orthorectified by a projective transform.

phenomenon in a broad range of systems, including polydisperse packings, consolidated soils and rocks, and more complex flows, such as multiphase, inertial, or non-Newtonian flows. While these cases may act to increase the rate of mixing due to the introduction of new flow phenomena, the fundamental kinematics described in this study must persist as they arise from the underlying topology of the grain pack. The investigation of these system-specific modulations of chaotic mixing in porous matter form promising research perspectives.

Materials and Methods

Experimental Protocol. The porous column consists of a vertically oriented rectangular column of cross-section 48×48 mm (Fig. 5), containing monodisperse borosilicate glass beads (Sigmund Lindner GmbH) of diameter d = 7, 10, or 20 mm that are loosely and randomly packed via gravitational settling. The pore space between the beads is filled with a glycerol-water mixture (1.4% wt/wt water) at controlled temperature (T = 25° C). Fluid flow is driven by a constant pressure difference imposed between the inlet and outlet ports, respectively, at the top and bottom of the column. The flow rate is continuously monitored at the outlet by a scale. Together with the knowledge of the packing porosity, this provides an estimate of the mean pore velocity, u. To visualize fluid flow and deformation through the bead pack, a solute fluorescent dye (PromoFluor-488LSS) is continuously injected in the upper part of the cell through a needle of internal diameter $L_0 = 0.5$ mm. The injected dye develops into a steady plume downstream of the injection point. The mean flow velocity u is chosen to be sufficiently low to get small Reynolds numbers and laminar flows (Re = $ud/\nu \approx 5 \cdot 10^{-3} \ll 1$, where $\nu \approx 700$ cP is the kinematic fluid viscosity), and fast enough for the dye Péclet number to be large and for fluid deformation to be measured from the dye backbone ($Pe = ud/D_m \approx 8.6 \cdot 10^3$, where $D_m \approx 2 \cdot 10^{-11} \text{ m}^2 \text{s}^{-1}$ is the molecular diffusivity of the dye in the glycerol-water mixture). For representativeness, experiments are repeated

for multiple dye-injection locations and for various packing realizations and bead diameters, as summarized in *SI Appendix*, Table S1. The coordinates of the bead centers are determined via a three-dimensional Hough transform on the image stack obtained by the translational laser scan, where the background fluid fluorescence allows distinguishing the grains. From these coordinates, several structural properties of the porous media are obtained: ϕ , the solid volume fraction (the ratio of volume occupied by the beads over the total column volume); z_c , the coordination number (the number of neighboring beads whose centers lie $d \pm 5\%$ away from the reference bead); and d_p , the mean diameter of the largest sphere inscribed in the pore space, obtained by a distance transform computed on a voxelized image of the fluid phase. **Reconstruction of the Dye-Filament Backbone.** In each cross-stream section

Reconstruction of the Dye-Filament Backbone. In each cross-stream section of the solute dye plume, a one-dimensional backbone of the dye filament is reconstructed via the adjustment of spline curves, and its total length *L* is computed (Figs. 2 and 6A). In experimental run II (Fig. 2 and Movie 51), the tracking is possible until downstream distance x = 9.27 d (beyond which the diffusive filament merges with itself), corresponding to a total elongation of $L/L_0 \approx 167$ (Fig. 2C), where L_0 is the initial length of the dye-filament backbone.

Distribution of Elongations. The sequential stretching and folding process leading to exponential growth of the total filament backbone length *L* implies that the length *I* of a fluid element follows a multiplicative random process (28, 32), such that *I* grows as $dI/dx^* = \gamma I$, where $x^* = x/d$ and γ is a random, statistically stationary stretching rate of mean λ and variance σ_{λ}^2 . The length *I* thus results from the product of the successive stretching rates γ , such that log *I* is the sum of a series of independent and identically distributed random variables. From the central limit theorem, the distribution of log *I* must then converge with downstream distance toward a normal distribution of mean $\lambda x'$ and variance $\sigma_{\lambda}^2 x^*$. This convergence is obtained after a few bead diameters since 1) the dye-filament lengths increase exponentially and thus sample an



Fig. 6. (*A*) Reconstruction of filament backbone (black line) from dye distribution and localization of high-curvature regions (red circles). (*B*) Local curvature along the filament backbone (black line) with isolated cusps (red circles) and threshold value $\kappa = 10^3 d^{-1}$ for cusp detection (dashed line). Note that the endpoints of filaments are considered as cusps, explaining why red circles can exist below κ . (*C*) Exponential growth of the number (n_c) of cusps in the filament backbone as a function of the advection distance (x/d) from the injection point. The fitted exponent for n_c (gray line) is similar to that for the total filament length (dashed line), suggesting stationarity of the stretching and folding processes.

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increasingly large number of independent stretching rates in the pore space and 2) the stretching rates are distributed within pores, so that several independent stretching rates are experienced over a unit diameter distance. The mean value of *I* is thus such that $\langle I \rangle \sim \exp(\mu x^*)$ with $\mu = \lambda + \sigma_{\lambda}^2/2$, which sets the growth rate of the total filament length to $\langle L/L_0 \rangle = \exp(\mu x^*)$.

Number of High-Curvature Regions. As shown in Fig. 1*B*, when the filament is advected through contact points between beads, localized regions of very high curvature develop in the filament backbone. We define cusps as isolated regions of the filament backbone where the curvature κ of the spline curve exceeds the threshold $\kappa = 10^3 d^{-1}$. We find (Fig. 6C) that the total number n_c of cusps in the filament backbone increases exponentially with the downstream distance x/d at a rate similar to that of the total filament length *L*. This suggests that stretching and folding events occur in proportion to each other and that the resulting process is statistically stationary.

Prediction of the Mean Stretching Rates from the Porous Media Properties. Based on the consistently observed sequence of stretching and folding in the pore space and its control by grain contacts (Fig. 1), we derive a general expression for the magnitude of the Lyapunov exponent in random granular media as a function of the coordination number z_c , the solid volume fraction ϕ_{c} the grain diameter d, and the mean pore diameter d_{p} . Since seqment lengths must double, on average, each time the segment encounters a contact, that is for each distance X_c , the Lyapunov exponent can be derived from Eq. 2 as $\lambda = d \log 2/X_c = d \log 2/(S_c/L_c)$, which requires estimating the mean segment length L_c and the mean surface area S_c swept by segments between successive contact points. A geometric estimate for the mean segment length L_c is obtained by assuming that an individual segment doubles its length at the constant exponential rate λ until it reaches the mean pore diameter d_p , where it inevitably collides with a contact point located at X_c downstream, e.g., $L_c(x) = d_p/2 \exp(\log 2 x/X_c)$. This expression yields the average value

$$L_c \approx \frac{d_p}{2\log 2}.$$
 [6]

From tomographic reconstruction of all of the experimental runs, the average pore diameter is $d_p \approx 0.24 d$ (Materials and Methods). Thus, we estimate $L_c \approx 0.17d$, which is in excellent agreement with the observations (Fig. 3). To develop an estimate of S_c , we consider the volume density ρ_V of contact points in the three-dimensional bead pack. As the number of grains per unit volume is $6\phi/(\pi d^3)$ and there are $z_c/2$ independent contacts per grain, the volume density of contact points is then $\rho_V = 3\phi z_c/(\pi d^3)$. Conversely, S_c is equal to the inverse of the areal density ρ_A of contact points in a filament sheet. To develop a relationship

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between ρ_V and ρ_A , we consider a series of filament sheets that arise from continuously injected line sources extending along length Z in the cross-stream z-coordinate of the column and are advected over distances X. These series of injection lines are also distributed along distance Y in the other cross-stream y-direction. If we consider the average number n_V of contact points in the volume V = XYZ, then the number n_A of contact points contained within a filament sheet is $n_A = \bar{\ell} / Y n_V$, where $\bar{\ell}$ is the average span of filament sheet in the y-direction that pass through a common contact point. The simplest estimate for $\bar{\ell}$ is obtained by considering the behavior of filament sheets in a open laminar flow over two spheres in contact at an angle α with the mean flow direction x. From symmetry arguments, the horizontal span $\ell(\alpha)$ is given by the projection in the plane normal to x of the contact line connecting the two sphere centers. Thus, $\ell(\alpha) = d \sin \alpha$, and $\overline{\ell}$ is given by the mean of $\ell(\alpha)$ over the distribution of contact angles. As $\rho_V \equiv n_V / (XYZ)$ and $\rho_A \equiv n_A / (XY)$, then $\rho_A = \bar{\ell} \rho_V$. For isotropic packings, contact lines are uniformly orientated in the space, and α is distributed as $p(\alpha) = \sin \alpha$. Averaging over this distribution then yields $\bar{\ell} \equiv \overline{d \sin \alpha} = d\pi/4$. From the results above, S_c for isotropic packings is then estimated as

$$S_{c} = 1/\rho_{A} = \frac{\pi d^{3}}{3\bar{\ell}\phi z_{c}}.$$
[7]

For anisotropic packings, the distribution of contact orientations may deviate from sin α , thus yielding different values for $\bar{\ell}$. Eqs. **6** and **7** provide the value of the mean fold distance $X_c = S_c/L_c$, and from Eq. **2**, the Lyapunov exponent reads

$$\lambda = \bar{\ell} \frac{3\phi z_c d_p}{2\pi d^2}.$$
 [8]

Using the estimates of ϕ , z_c , and d_p obtained from tomographic reconstruction and $\bar{\ell}/d = \pi/4 \approx 0.78$ for isotropic packings, Eq. 8 yields $\lambda = 0.27$, a value comparable to the experimental estimate ($\lambda = 0.21$). The slight overestimate of λ can be explained by the anisotropy introduced by the gravitational packing of the beads and the finite size of the experimental column. Indeed, independent measurements of $\rho_V = 2.86 \ d^{-3}$ and of $S_c = 0.58 \ d^2$ (Fig. 3) indicate that, experimentally, $\bar{\ell}/d = (d\rho_V S_c)^{-1} \approx 0.6$, instead of $\bar{\ell}/d = 0.78$ expected for isotropic packings. Using this value in Eq. 8 yields $\lambda \approx 0.21$, in much better agreement with the experimental estimate.

Data Availability. All data needed to evaluate the conclusions in this paper are available in the main text or in *SI Appendix*.

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