The infiltration of a solute in a fractal porous medium is usually anomalous, but chemical reactions of the solute and that material may increase the porosity and affect the evolution of the infiltration. We study this problem in two- and three-dimensional lattices with randomly distributed porous sites at the critical percolation thresholds and with a border in contact with a reservoir of an aggressive solute. The solute infiltrates that medium by diffusion and the reactions with the impermeable sites produce new porous sites with a probability \( r \), which is proportional to the ratio of reaction and diffusion rates at the scale of a lattice site. Numerical simulations for \( r \ll 1 \) show initial subdiffusive scaling and long time Fickean scaling of the infiltrated volumes or areas, but with an intermediate regime with time increasing rates of infiltration and reaction. The anomalous exponent of the initial regime agrees with a relation previously applied to infinitely ramified fractals. We develop a scaling approach that explains the subsequent time increase of the infiltration rate, the dependence of this rate on \( r \), and the crossover to the Fickean regime. The exponents of the scaling relations depend on the fractal dimensions of the critical percolation clusters and on the dimensions of random walks in those clusters. The time increase of the reaction rate is also justified by that reasoning. As \( r \) decreases, there is an increase in the number of time decades of the intermediate regime, which suggests that the time increasing rates are more likely to be observed is slowly reacting systems.

\[ I \sim t^n, \]

where \( n \) is termed infiltration exponent. If the medium in \( d \) dimensions is homogeneous and the source is a border of dimension \( d-1 \), the infiltration is Fickean, with \( n = 1/2 \), as shown in models and experiments [5, 8]. However, anomalous exponents \( n \neq 1/2 \) were already obtained in studies of moisture infiltration in construction materials [9, 13], in several models of infiltration in regular fractals [5, 6, 14], and in the infiltration of glycerin in Hele-Shaw cells with fractal pore geometry (a problem that is equivalent to diffusive infiltration) [7].

These processes are related to the molecular diffusion in the medium. When particles randomly move in a homogeneous medium, their root-mean-square displacement \( R \) increases in time as \( t^{1/2} \) (i.e. Fickean diffusion). In highly disordered media, anomalous diffusion is observed, in which

\[ R \sim t^{1/d_W}, \]

with the random walk dimension \( d_W \neq 2 \) [15, 18]. In fractal porous media, the self-similar distributions of irregularities, such as impenetrable barriers or dead ends, usually lead to subdiffusion, in which \( d_W > 2 \). However, note that the infiltration exponent \( n \) and the random walk exponent \( 1/d_W \) may be different. For instance, in infinitely ramified fractals such as Sierpinski carpets and Menger sponges, the connection of diffusion driven infiltration and the diffusion anomalies led to a relation between \( n, d_W \), the fractal dimension \( d_F \) of the medium, and the dimension \( d_B \) of the infiltration border [5, 8]. A possible consequence is that superdiffusive infiltration \((1/2 < n \leq 1)\) occurs in a medium where random walks are subdiffusive [14].

The interplay of infiltration in disordered media and chemical reactions may lead to changes in the structure of those media, which is of particular importance in the evolution of geological materials [19, 22]. Diffusion is expected to be the dominant transport mechanism in several cases, particularly in low porosity media [1, 23, 24]. For instance, in olivine-rich rocks whose porosity is near a few percent, serpentinization creates small fractures that serve as pathways for fluid infiltration [20, 25]. Moreover, water infiltration and loss of reaction products in basalt clasts lead to the formation of porous weathered rinds around (more compact) unaltered cores [1, 25]. Fractal pore geometry was confirmed in both altered and unaltered domains of those clasts [26] and was also observed in many other geological materials [30, 31]. Infiltration

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I. INTRODUCTION

When a porous medium is filled with a static fluid and an external surface is put in contact with a reservoir of a mobile species with a different concentration, that species may infiltrate into or out of that medium by diffusion. This occurs, for instance, in the weathering of geological materials [1], in the dispersion of contaminants in soils [2], and in the transport of radionuclides in nuclear waste containers [3]. A simple model for the diffusive infiltration without reactions considers lattice random walks of (excluded volume) particles that start from a source at one border, as originally proposed by Sapoval et al [4] and recently extended to fractal geometries [5]. The infiltration length \( I \) is defined as the infiltrated volume per unit area of the exposed surface and is shown to scale as

\[ I \sim t^n, \]

where \( n \) is termed infiltration exponent. If the medium in \( d \) dimensions is homogeneous and the source is a border of dimension \( d-1 \), the infiltration is Fickean, with \( n = 1/2 \), as shown in models and experiments [5, 8]. However, anomalous exponents \( n \neq 1/2 \) were already obtained in studies of moisture infiltration in construction materials [9, 13], in several models of infiltration in regular fractals [5, 6, 14], and in the infiltration of glycerin in Hele-Shaw cells with fractal pore geometry (a problem that is equivalent to diffusive infiltration) [7].

These processes are related to the molecular diffusion in the medium. When particles randomly move in a homogeneous medium, their root-mean-square displacement \( R \) increases in time as \( t^{1/2} \) (i.e. Fickean diffusion). In highly disordered media, anomalous diffusion is observed, in which

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anomalies are expected due to the fractality, but they may change with the progress of the reactions. Such changes were recently shown in a model of infiltration and reaction in infinitely ramified fractals, in which the initial subdiffusive behavior crossed over to a long time Fickian infiltration \[32\].

This scenario motivates the present investigation of the coupling of diffusive infiltration and dissolution reactions (which create new porosity) in critical percolation clusters \[31, 33\], which are the most prominent stochastic fractal media exhibiting anomalous diffusion. Our first step is to study the non-reactive infiltration of a solute in hypercubic lattices of dimensions \(d = 2\) and \(3\) where the porous sites are randomly distributed with the critical percolation probabilities and the remaining sites are impermeable. The infiltrated area \((d = 2)\) and volume \((d = 3)\) are shown to increase with exponents \(n < 1/2\) consistent with the same scaling relation obtained in infinitely ramified fractals \[3\]. The second part of this work considers that reactions between the infiltrating solute and the impermeable solid form new porous sites in which solute transport is possible. The initial subdiffusive infiltration and the long time Fickian behavior are observed, but they are separated by a regime in which the infiltrated and the dissolved volumes (or areas) increase faster than linearly in time, i.e. infiltration and dissolution have time increasing rates. This phenomenon results from the combination of a slow penetration of the solute in a preferential direction of a porous system of vanishing density and the linear advance in all directions of the dissolution of the solid walls surrounding the pores. The time range of this regime is shown to increase as the reaction rate decreases (relatively to the diffusion rate), so the nontrivial anomaly may last for long times in slowly reacting materials with fractal pore systems. These results are obtained with numerical simulations and explained by a scaling approach.

This paper is organized as follows. Sec. II presents the model of infiltration and reaction, information on the methods of solution, and the main quantities to be measured. Sec. III reviews the results for the same model with an initially compact (not porous) medium, which helps the interpretation of some results in the porous media. Sec. IV presents results for infiltration without reactions in \(d = 2\) and \(3\). Sec. V presents simulation results for the model with reactions. Sec. VI presents a scaling approach that explains the observed evolution of infiltration and reaction lengths, with particular emphasis on the regime with time increasing rates. Sec. VII summarizes our results and presents our conclusions.

II. MODELS AND METHODS

A. Model Definition

The porous media are built in the region \(z > 0\) of hypercubic lattices of dimension \(d\). Each site is expected to represent a homogeneous mesoscopic region of a porous material. The permeable sites, which are termed P sites, are inert. Their initial fraction is equal to the percolation threshold \(p_c\), with nearest neighbor (NN) connectivity. The impermeable sites contain a reactive material and are termed M sites; their initial fraction is \(1 - p_c\). With these definitions, solute molecules can be transported only through P sites that are NN.

The pore solution and the material M are initially in chemical equilibrium. A large solution at \(z \leq 0\) with a constant concentration of an aggressive solute is put in contact with the medium at time \(t = 0\). Thus, \(z = 0\) is the infiltration border, whose dimension is \(d_B = d - 1\). The solute is represented by S particles that permanently fill all the sites with \(z = 0\) and that may also occupy P sites of the medium. Excluded volume conditions are considered for the S particles, so that a P site or a border site may have zero or one S particle. The possible configurations of the lattice sites are shown in Fig. 1(a). Figure 1(b) shows a configuration of a lattice in \(d = 2\) after infiltration of some S particles and the filled border at \(z = 0\).

In a time interval \(\tau\), all S particles attempt to hop to a randomly chosen NN site. If an S particle attempts to hop from the border \(z = 0\) to an empty P site, the hop is executed and another S particle is immediately inserted at its initial position, so that the border remains with the same solute concentration; this is illustrated in Fig. 1(c). However, if the S particle attempts to hop to an M site or to a site with another S particle, the attempt is rejected and that particle remains in the same position. Fig. 1(d) illustrates the hop of an S particle at a P site, which was executed because the target site was an empty P site; however, if that particle attempted to hop to the site below it or to the site above it, the attempt would be rejected.

In the same time interval \(\tau\), an S particle in contact with a NN M site can react with probability \(r\). The reaction leads to the transformation of the M site into an empty P site and to the annihilation of the S particle, as illustrated in Fig. 1(e). This rule of the model is a simplified description of a series of physical and chemical processes: (i) the reaction between S and M forms soluble and non-soluble products; (ii) the non-soluble products form a porous precipitate with a physical structure similar to those of the initial P sites; (iii) chemical equilibrium is restored in the pore solution where the reaction occurs (this is represented by the annihilation of the S particle).

B. Interpretation of the model parameters

Here we partly follow the reasoning of Ref. \[32\] to relate the model parameters in \(d = 3\) to measurable quantities.

Letting \(a\) be the lattice constant and \(D\) be the diffusion coefficient of the S particles in the porous medium formed
only by P sites, we have

\[
D = \frac{a^2}{2d\tau}.
\]

This coefficient is expected to be smaller than that in free solution.

When an M site reacts, the change in the number of moles of the reacting material is \(\Delta n_M = a^2/v\), where \(v\) is the molar volume of that material; in an application, \(v\) depends on the properties of the reacting material and its volume fraction in the solid represented by the M site. The area of the M site in contact with a NN P site is \(a^2\), but the P site represents a mesoscopic porous region, so only a fraction of that area is in contact with pore solution. We assume that this fraction is equal to the effective porosity of a P site, which is denoted as \(\phi_P\); see e.g. Ref. [34]. Thus, an area \(A_M = \phi_P a^2\) of the M site is in contact with the solution in the NN P site. Since the \(\Delta n_M\) moles of M react with a probability \(r\) in a time interval \(\tau\), the reaction rate \(k\), in mol/(m\(^2\)s), is given by

\[
k = r\frac{\Delta n_M}{(A_M\tau)} = 6rD/(av\phi_P);
\]

here Eq. (3) was used with \(d = 3\). This gives

\[
r = \frac{\phi_Pv a k}{6D}.
\]

Thus, \(r\) is a ratio between rates of reaction and diffusion in the volume of a lattice site, which may be interpreted as a second Damkohler number of the model [35].

The condition of slow reaction compared to diffusion implies \(r \ll 1\), which is considered throughout this work. Since our main results are obtained in terms of the parameter \(r\), the application of the model to a real system is possible if the physical and chemical parameters in Eq. (4) are estimated.

C. Methods of Solution

Scaling relations are well known for the structural properties of percolation clusters and the diffusion of tracers in those media [16, 33, 36]. Relations between infiltration and diffusion exponents in regular fractals are also known [5, 8] and can be tested in the infiltration model without reactions at \(p = p_c\). Moreover, scaling approaches for the interplay of diffusion and alteration reactions (interfacial dissolution reprecipitation mechanism [37]) were developed in Ref. [38] for non-porous media; an analogous approach was used in the study of the growth of passive layers on metallic surfaces [39]. A combination of these methods is used here to explain the scaling regimes of infiltration and reaction.

We also use kinetic Monte Carlo simulations to support the predictions of this scaling approach. In \(d = 2\), we consider lattices with infiltration border \((z = 0)\) of size \(L = 1024a\), with very large length in the \(z\) direction, and with periodic boundary conditions in the other direction. Some simulations are also performed in \(L = 2048a\) to discard the possibility of finite-size effects. The initial fraction of pore sites is \(p_c = 0\), the values of \(r\) range from \(10^{-2}\) to \(10^{-6}\), and 100 realizations are used to calculate average quantities, up to the maximal simulation time \(10^7\tau\). In \(d = 3\), we consider lattices with infiltration border of size \(L = 256a\), with very large length in the \(z\) direction, and with periodic boundary conditions in the other directions. Some simulations are also performed in \(L = 512a\) to discard the possibility of finite-size effects. The initial fraction of pore sites is \(p_c = 0\), the values of \(r\) range from \(10^{-2}\) to \(10^{-5}\), and 50–100 realizations are used to calculate average quantities. The maximal simulation time is \(10^5\tau\) for most samples, but \(10^6\tau\) for the smallest \(r\).
A medium at $p_c$ has a percolating cluster of $P$ sites with fractal dimension $d_F = 91/48$ in $d = 2$ \cite{12,13} and $d_F = 2.5230 \pm 0.0001$ in $d = 3$ \cite{14}. The random walk dimension in critical percolation clusters in $d = 2$ obtained from simulations is $d_W = 2.8784 \pm 0.0008$ \cite{15}. In $d = 3$, the scaling of the conductivity in percolation clusters gives $d_W = 3.806 \pm 0.001$ \cite{14,46,47}. These values are used to test the scaling relations determined here.

D. Basic Quantities

The infiltration time is denoted as $t$ and a dimensionless infiltration time is defined as

$$ T = \frac{t}{\tau}. \quad (5) $$

The number of infiltrated sites, i.e., $P$ sites with a particle $S$, is denoted as $N_I$. For a unified description in all spatial dimensions, we define a dimensionless infiltration length $I$ as the ratio between the number of infiltrated sites, $N_I$, and the number of sites of the infiltration border, $(L/a)^d$:

$$ I = \frac{N_I}{(L/a)^d}. \quad (6) $$

From this definition, $Ia$ can be interpreted as the average length in the $z$ direction occupied by the (non-contiguous) infiltrated $P$ sites. The part of the medium with distance $\leq Ia$ from the infiltration border is termed the infiltrated region; it comprises most sites with $S$ particles.

The number of $M$ sites transformed into $P$ sites (i.e., $M$ sites that react) is denoted as $N_R$. The dimensionless reaction length $R$ is defined as the ratio between this number and the number of sites of the infiltration border, $(L/a)^d$:

$$ R = \frac{N_R}{(L/a)^d}. \quad (7) $$

The length $Ra$ measures the extent of penetration of the reaction in the $z$ direction.

III. REVIEW OF THE MODEL WITH COMPACT REACTIVE MEDIA

Here we consider the case in which the medium interacting with the solution is compact, i.e., initially all lattice sites with $z > 0$ are $M$ sites. Figure 2 shows snapshots of the attacked solid and the infiltrated region at several times for $r = 10^{-2}$.

In the shortest times, we observe a large density of $S$ particles near the $M$ surface because they can rapidly move in the porous medium near the source, whereas their reactions with $M$ sites are relatively slow. Thus, the $M$ sites at the surface are almost all the time in contact with an $S$ particle (e.g., in the first three panels of Fig. 2). This is a regime controlled by the reaction, in which $I \sim rT$, as shown in Fig. 3. However, when the average distance between the source and the $M$ surface becomes large, the time for a new particle to leave the source and to reach that surface increases. This time will eventually be larger than the average time $(r^{-1})$ for a reaction with an $M$ site, which leads to a depletion in the density of $S$ near the $M$ surface. In this condition, the infiltration is controlled by the diffusion of the $S$ particles across the length $I$, so $I \sim T^{1/2}$ ($R$ exceeds $I$ because the $S$ particles occupy only a fraction of the sites that reacted). This is the long time Fickean regime, as shown in Fig. 3.

The crossover time between the reactive and the Fickean regime is obtained by matching the expressions for the infiltration length in those regimes:

$$ T_c \sim r^{-2}. \quad (8) $$

The right panel of Fig. 2, which is at $T \approx T_c \sim 10^4$, actually shows that the density of $S$ particles near the $M$ surface is much smaller than the density at the shortest times. Fig. 3 confirms that the infiltration is entering the Fickean regime at that time.

For further comparison, note that the crossover shown in Fig. 3 occurs with a continuous decrease of the slope of the log $I \times \log T$ and $\log R \times \log T$ plots.

IV. INFILTRATION WITHOUT REACTIONS

Figure 4 shows snapshots of an infiltrated medium in $d = 2$ at several times. The anomalous advance of the infiltration front is confirmed by the infiltration length bars: as the time increases by a factor 4 between consecutive panels, $I$ varies by factors smaller than 2 (the factor 2 is expected in Fickean infiltration). In $d = 3$, $I$ varies by smaller factors when results at the same time are compared. As the infiltration advances, we also observe a smaller density of filled pore branches, which is related to the fractality of the percolation cluster.

In Ref. [5], a scaling approach predicted that the infiltration length in fractal porous media scales as in Eq. (1) with

$$ n = \frac{d_F - d_B}{d_W}. \quad (9) $$

Eq. (9) was confirmed in several infinitely ramified fractals, including some cases with superdiffusive infiltration $(1/2 < n < 1)$ \cite{8,14}. However, it was not formerly tested in stochastic fractals.

Using the structural and dynamical exponents of percolation, Eq. (9) predicts $n \approx 0.3113$ in $d = 2$ and $n \approx 0.137$ in $d = 3$. Figs. 5a and 5b show the evolutions of the infiltration length in percolating media in $d = 2$ and $d = 3$, respectively, with dashed lines indicating the predicted exponents $n$. The agreement in $d = 2$
FIG. 2. (Color online) Infiltration in a two dimensional compact medium with $r = 10^{-2}$. The left (green) bar and the right (blue) bar represent the infiltration and reaction lengths, respectively.

FIG. 3. (Color online) Evolution of the infiltration and reaction lengths with an initially compact medium in $d = 2$ and $r = 10^{-2}$.

is very good, but deviations are observed in $d = 3$ until the longest simulated times. The local slopes of the log $I \times \log T$ plots are the effective infiltration exponents $n_{\text{eff}}$ shown in the insets of Figs. 5(a) and 5(b). In $d = 2$, $n_{\text{eff}}$ reaches values very close to the prediction of Eq. (9) at $T \sim 10^5$. In $d = 3$, $n_{\text{eff}}$ is more than 10% larger than the predicted value at $T \sim 10^6$. The fit of $n_{\text{eff}}$ as a function of $T^{-1/6}$ leads to an asymptotic estimate consistent with that of Eq. 9, which shows the presence of large corrections to the dominant scaling.

Note that the right panel in Fig. 4 shows that the distances of several S particles from the source are much larger than $I$. This occurs because their displacement is governed by Eq. (2), with exponent $1/d_W$, while $I$ increases with a smaller exponent $n$ [Eq. 9] with $d_F - d_B < 1$ implies $n < 1/d_W$.

V. SIMULATIONS OF INFILTRATION WITH REACTIONS

A. Results in $d = 2$

Figure 6 illustrates the infiltration in $d = 2$ for $r = 10^{-3}$ and the same times of Fig. 4 (the case without reactions). The results for $T < 10^5$ show a slow advance of the infiltration front because the reaction probability $\sim rT$ is small; this is similar to the case without reactions. However, at longer times, the S particles fill some dissolved (or partly dissolved) layers near the source. The advance of the infiltration is faster than that without reactions. The reaction length $R$ increases even faster. If the results at $T = 9600$ are compared with the case of a compact medium (Fig. 2), here we observe a deeper infiltration despite the smaller value of $r$. This is possible because the solute infiltrates into the channels of the porous medium and dissolves the channel walls, which facilitates the subsequent infiltration.

Figures 7(a) and 7(b) show the time evolution of the infiltration length and of the reaction length, respectively, for several values of $r$. At short times, the scaling of $I$ is subdiffusive, similarly to the case without reactions; in this regime, $R$ is very small, but rapidly increases in time. The extent of the reactions eventually increase and lead to deviations from the subdiffusive scaling. The time for these deviations to appear increases as $r$ decreases; inspection of Figs. 7(a) and 7(b) shows that they occur as $R \sim 1$ in all cases. Subsequently, rapid increases of $I$ and $R$ are observed, followed by the convergence of both quantities to the Fickian behavior $\sim T^{1/2}$ (the same
Figures 4 and 5 show the evolution of the infiltration rate $I \equiv dI/dT$ and reaction rate $R \equiv dR/dT$, respectively. $I$ decreases at short times, has a local minimum just after the subdiffusive regime, and has a local maximum just before the Fickian regime. Between those extrema, $I$ slowly increases in time; the time interval of this intermediate regime increases as $r$ decreases. $R$ increases as a power law in the subdiffusive regime of $I$ and subsequently shows a faster time increase (which can hardly be fit as a power law); at longer times, it also shows a maximum before the crossover to the Fickian regime.

The crossover time between the subdiffusive infiltration and this intermediate regime is denoted as $T_1$ and the crossover time to the Fickian regime is denoted as $T_2$. Here we consider two possible definitions of $T_1$: in the first one, it is the time of the local minimum of $I$ after the subdiffusive scaling; in the second one, it is the time in which the relative deviation of $I$ from the non-reactive case ($r = 0$) reaches a preset value $\Delta$ (this is the same definition used in Ref. [32] for a similar model in regular fractals). The crossover time $T_2$ is also estimated considering two definitions: in the first one, it is the time of the local maximum of $I$ before the Fickian regime; in the second one, it is the time in which the relative deviation of $I$ from the asymptotic Fickian scaling reaches a preset value $\Delta$. The latter considers the exact asymptotic relation $I = 2\sqrt{Dt/(d\pi)}$ for a lattice with only P sites.

Figures 7(c) and 7(d) show the evolution of the infiltration rate $\dot{I}$ and reaction rate $\dot{R}$, respectively. $\dot{I}$ decreases at short times, has a local minimum just after the subdiffusive regime, and has a local maximum just before the Fickian regime. Between those extrema, $\dot{I}$ slowly increases in time; the time interval of this intermediate regime increases as $r$ decreases. $\dot{R}$ increases as a power law in the subdiffusive regime of $I$ and subsequently shows a faster time increase (which can hardly be fit as a power law); at longer times, it also shows a maximum before the crossover to the Fickian regime.

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Figures 8(a) shows the crossover times calculated with the two definitions as a function of $r^{-1}$. In this analysis, we used $\Delta = 20\%$. The plots suggest power law scalings as

\[ T_1 \sim r^{-b}, \quad T_2 \sim r^{-c}, \quad (10) \]

where $b$ and $c$ are positive exponents. Linear fits of the $T_1$ data for $r \leq 10^{-3}$ give $b = 0.79$ with the two methods;
the fits of the $T_2$ data give $c = 1.03$ and 1.08. Since $c$ is unequivocally larger than $b$, the ratio $T_2/T_1$ increases as $r$ decreases, i.e. the time interval of the intermediate regime increases. It may be very long for slow reactions; for instance, for $r = 10^{-6}$, $T_2$ and $T_1$ differ by approximately 1.5 order of magnitude.

The values of $I$ at $T_1$ and $T_2$, which we denote as $I_1$ and $I_2$, respectively, are shown in Fig. 8(b) as a function of $r^{-1}$. As $r$ decreases, $I_1$ increases, which is consistent with a longer subdiffusive regime; $I_2$ increases slightly faster, which means that a relatively larger infiltration is obtained in the regime with time increasing rates. The values of $R$ at those crossovers, which we denote as $R_1$ and $R_2$, respectively, are shown in Fig. 8(c). $R_1 \sim 1$ for all values of $r$, which suggests that this condition is related to the breakdown of the subdiffusion. In Fig. 8(d), we show the infiltration rates $I_1$ and $I_2$ at the crossovers, and in Fig. 8(e) we show the corresponding reaction rates $R_1$ and $R_2$. In all cases, a decrease of these rates is observed as $r$ decreases (note that they are rates calculated at different times, so it does not contradict the observation of time increasing rates in the intermediate regime for a constant $r$). In Figs. 8(b)-(e), the linear fits show that those quantities vary as power laws of $r$ for small values of this parameter; the exponents shown in the plots weakly depend on the definition used to calculate the crossover times.

**B. Results in $d = 3$**

Fig. 9 shows cross sections of a medium during its infiltration with reaction probability $r = 10^{-4}$. In the first two panels, the infiltration is slow and a small number of M sites has reacted. The third panel shows a high filling of the accessible region near the source, but with no significant change due to the reaction. Between the third and the fourth panel, the infiltration and the reaction rapidly advance. This is accompanied by significant dissolution of some layers near the source (within the reaction length), where the density of S particles is high. The layers within or slightly below the infiltration length are also highly filled, but only a fraction of the M sites have reacted.

Figures 10(a) and 10(b) show the evolution of $I$ and $R$, respectively, for several $r$, and Figs. 10(c) and 10(d) show the corresponding rates. The qualitative evolution is similar to that in $d = 2$, with an initial subdiffusive regime, an intermediate regime with time increasing rates, and a final Fickean regime. However, the distinct features of the intermediate regime are sharper in $d = 3$: the time increase of $I$ is clearer and the increase of $R$ is faster in comparison with $d = 2$. Moreover, this regime lasts longer as $r$ decreases.

Here we also characterize the crossovers between the three scaling regimes by the times $T_1$ and $T_2$, using the two definitions given in Sec. VA. For the method that considers fractional deviations of $I$ from the subdiffusive and from the Fickean scalings, here we use $\Delta = 50\%$. The crossover times are shown in Fig. 11(a) as a function of

**FIG. 6.** (Color online) Infiltration with dissolution ($r = 10^{-3}$) in a porous medium at the percolation threshold in $d = 2$. In each panel, the left green bar represents the infiltration length and the right blue bar represents the reaction length.
FIG. 7. Evolution of (a) infiltration length and (b) reaction length in $d = 2$ for several values of $r$; the dashed lines are theoretical predictions for the subdiffusive and the Fickian regime. (c) and (d) shows the respective rates of change, with dashed lines indicating the slopes theoretically predicted for the subdiffusive and the intermediate regimes.

$r^{-1}$ and also follow the power law relations of Eq. [10]. The exponents for $T_1$ are $b = 0.77$ and $0.81$, while the exponents for $T_2$ are $c = 1.05$ and $1.10$; thus, the different definitions of those times also lead to small differences in the measured exponents. Since $c > b$, the number of

FIG. 8. (Color online) (a) Crossover times $T_1$ and $T_2$ as a function of $r^{-1}$, in $d = 2$, obtained from local extrema of $I$ (labels $\text{min}$ and $\text{max}$) and from the deviation $\Delta = 20\%$. (b) Infiltration lengths ($I_1, I_2$) and (c) reaction lengths ($R_1, R_2$) at the crossover times [same color code as (a)]. (d) Infiltration rates ($I_1, I_2$) and (e) reaction rates ($R_1, R_2$) at the crossover times [same color code as (a)]. In all plots, the error bars are smaller than the sizes of the symbols and the slopes of the least squares fits of all data sets (dashed lines) are indicated.
FIG. 9. Evolution of a small cross-section of the lattice in $d = 3$ with $r = 10^{-4}$. The time 625 is below $T_1$, 2500 is close to $T_1$, 10000 is close to the inflection point, and 40000 is close to $T_2$. The left green bar represents the infiltration length and the right blue bar the reaction length.

time decades of the intermediate regime $[\log_{10}(T_2/T_1)]$ increases as $r$ decreases.

The infiltration lengths at $T_1$ and $T_2$ are shown in Figure 11(b) as a function of $r^{-1}$; the reaction lengths at the crossover times are shown in Fig. 11(c). As in the two-dimensional case, here we observe that $I_1$ increases as $r$ decreases, consistently with a longer subdiffusive regime, whereas $I_2$ increases faster, consistently with larger infiltrations during the intermediate regime of time increasing rates. We also observe that $R_1 \sim 1$ for all $r$, supporting the assumption that this condition is related to the breakdown of the subdiffusion. Figure 11(d) shows the infiltration rates and Fig. 11(e) shows the reaction rates at the crossover times $T_1$ and $T_2$, which follow similar trends as the crossover rates in $d = 2$. In Figs. 11(b)-(e), the scaling exponents obtained from data fits with the smallest $r$ are shown in the plots and are weakly dependent on the definition of the crossover times.

VI. SCALING APPROACH

For any reaction probability $r$, the short time subdiffusive infiltration is described by Eq. (9) because the number of dissolved M sites is small. Reactions may occur on the walls of the infiltrated channels, i.e. on the faces of the M sites in contact with S particles. An analogy with the dissolution of the compact medium (Sec. III) is helpful here: at short times, Figs. 2(a)-(c) show that almost all P sites in contact with the M sites are infiltrated. Thus, the dissolved region acts like an extension of the source at times much shorter than the crossover to the Fickian behavior. Here, the difference is that the S particles also invade the original P sites of the porous medium, i.e. the extended source is inside the porous medium.

This reasoning implies that the number of faces of M sites in contact with S particles is proportional to the number $N_I$ of infiltrated P sites. In $d = 3$, the area of M sites in contact with S particles is of order $A_I \sim N_I a^2 \sim IL^2$; in $d = 2$, the length in contact with S particles is $L_I \sim N_I a \sim IL$. These relations omit the average number of faces of M sites per infiltrated P site, which is of order 1.

In the neighborhood of an infiltrated P site, an M site reacts with probability $r$ in a time $\tau$. Thus, in a time $t$, the reaction advances into an average length $l_\perp \sim a(\tau/r)t = arT$ of the M sites surrounding that P site. This is applicable in $d = 2$ and $d = 3$ at short times, in which $l_\perp \leq a$, i.e. in which the advance of the reaction is limited to the M sites in the closest neighborhood of the P site. The total volume of M that reacts in a time $T$ is $V_R \sim A_I l_\perp \sim a L^d r T$ in $d = 3$; in $d = 2$, the total area that reacts is $A_R \sim L_I l_\perp \sim a L r T$. Using Eqs. 7
and (9), the reaction length in both dimensions scales as

\[ R \sim rT^{n+1}. \]  

(11)

Since \( n > 0 \), \( R \) increases faster than linearly since short
times. The corresponding rate of reaction is

\[ \frac{dR}{dt} \sim rT^n. \tag{12} \]

In \( d = 2 \), the plots in Fig. 7(d) have slopes consistent with the estimate \( n \approx 0.3113 \). In \( d = 3 \), the plots in Fig. 10(d) have slopes slightly larger than the estimate \( n \approx 0.137 \); however, this is expected because the effective exponents \( n_{\text{eff}} \) are large at short times, as shown in the inset of Fig. 5(b).

After some time, several M sites are transformed into P sites, which facilitates the infiltration of other S particles. Figs. 8(c) and 11(c) show that the first crossover occurs when \( R \sim 1 \), which corresponds to the reaction of one M site per source site. This means that the extended source formed inside the porous medium has a volume of the same order as the volume of the source localized at the infiltration border. After it occurs, the attack to the M sites inside the medium (by an increasing number of S particles) is faster than the attack to the top M sites; the subdiffusive behavior then ceases and a different scaling regime begins. The crossover time \( T_1 \) is obtained by substituting the condition \( R \sim 1 \) in Eq. (11), which leads to the scaling in Eq. (10) with exponent

\[ b = \frac{1}{1 + n}. \tag{13} \]

The infiltration and reaction lengths at \( T_1 \) are obtained by substituting Eqs. (10) and (13) in Eqs. (9) and (11); their time derivatives give the rates of those lengths at the crossover:

\[ I_1 \sim r^{-n/(n+1)} \quad , \quad \dot{I}_1 \sim r^{(1-n)/(1+n)} \quad , \quad \dot{R}_1 \sim r^b. \tag{14} \]

Our numerical results support these scaling relations. In \( d = 2 \), the estimate \( n \approx 0.3113 \) gives \( b \approx 0.763 \), \( n/(n+1) \approx 0.237 \), and \( (1-n)/(1+n) \approx 0.525 \) for the exponents in Eq. (14). The numerical values of the exponents of \( T_1 \), \( I_1 \), \( T_1 \), and \( \dot{R}_1 \), shown in Figs. 8(a)-(c), are close to the predictions of this scaling approach, with maximal deviations \( \approx 6\% \). In \( d = 3 \), the asymptotic value \( n \approx 0.137 \) gives \( b \approx 0.880 \), \( n/(n+1) \approx 0.120 \), and \( (1-n)/(1+n) \approx 0.759 \). The numerical estimates of the exponents of \( I_1 \), \( I_1 \), and \( \dot{R}_1 \) [Figs. 11(a)-(c)] differ up to \( \approx 13\% \) from those values. The numerical estimate of the exponent of \( I_1 \) [Fig. 11(b)] has a larger deviation. In this case, \( T_1 \) is in the range \( 10^2-10^6 \) [Fig. 11(a)], in which the infiltration length without reactions increase with \( n_{\text{eff}} \) between 0.25 and 0.18 [Fig. 5(b)], which are 30\%–80\% larger than the asymptotic value \( n \approx 0.137 \). Thus, the numerical estimates of exponents \( b \) and \( (1-n)/(1+n) \) are actually expected to be smaller than the scaling predictions based on the asymptotic \( n \), whereas the estimate of \( n/(n+1) \) is expected to be larger than the scaling prediction.

The deviation from subdiffusion at \( T_1 \) implies that the fractality is broken in the region within the infiltration length. At \( T > T_1 \), the dissolution of M sites creates new paths for the infiltration. The S particles fill P sites that belong to the critical percolation cluster plus new P sites created by the reactions and P sites that were initially isolated; see Fig. 12. To estimate the infiltration length, we consider these contributions separately.

FIG. 12. (Color online) Scheme of an infiltrated region of the porous medium in \( d = 2 \) before (left) and after (right) the reactions occur.

First consider the infiltrated part of the original percolation cluster. The infiltration source is extended, i.e. the non-infiltrated parts of that cluster are below a region with a high density of S particles. Thus, the infiltration advance is expected to be the same observed when the initial percolation cluster was in contact with the source at \( z = 0 \). In each time interval \( T_1 \), the infiltration advances into that cluster by a length \( I_1 \) in the \( z \) direction (as in the initial subdiffusive regime), which means that the extended source advances by that length. At \( T > T_1 \), the infiltration length \( I_{\text{pc}} \) in the original percolation cluster is proportional to \( T/T_1 \):

\[ I_{\text{pc}} \sim I_1 \frac{T}{T_1}. \tag{15} \]

Now we analyze the infiltration of the other P sites, i.e. sites that do not belong to the original percolation cluster. The reactions around an infiltrated P site advance in all directions to a length \( l_\perp \sim a (r/rT) t = arT \), as illustrated in Fig. 12; now \( l_\perp \) is not of the same order as \( a \). In a \( d \)-dimensional region with size \( l_\perp \), the total number of P sites is \( \sim p_s (l_\perp/a)^d \), i.e. the P sites occupy a finite fraction of the available volume. In the same region, the number of P sites in the critical percolation cluster is \( \sim (l_\perp/a)^d_F \). Thus, the ratio between the total number of P sites and the number of P sites in the percolation cluster at time \( T \) is

\[ f_p (T) \sim \left( \frac{l_\perp}{a} \right)^{d-d_F} \sim (rT)^{d-d_F}. \tag{16} \]

Since \( d > d_F \), this result implies that most of the infiltrated sites are not those of the original percolation cluster.

Recalling that \( I_{\text{pc}} \) [Eq. (15)] accounts only for the infiltrated sites of the percolation cluster, the total infiltrated
length $I$ is larger than $I_{pc}$ by the factor $f_p(T)/f_p(T_1)$:

$$I \sim \frac{f_p(T)}{f_p(T_1)} f_{pc} \sim r^h t^g$$

with $g \equiv 1 + d - d_F$, $h \equiv d - d_F$.

This gives $I \sim I_1$ at $T \sim T_1$. The infiltration rate scales as

$$\dot{I} \sim r^h T^{d-d_F}.$$  \hspace{1cm} (17)

Eqs. (17) and (18) give $g > 1$ in any dimension, so they theoretically predict the time increasing infiltration rate. The numerical values are $g = 53/48 \approx 1.104$ in $d = 2$ and $g \approx 1.477$ in $d = 3$, which show that the effect is more pronounced in $d = 3$. This result is confirmed with good accuracy by the slopes $d - d_F \approx 0.104$ and 0.477 of the infiltration rate evolution in Figs. 7(c) and 10(c), respectively.

The crossover to Fickean infiltration occurs when the infiltration length of Eq. (17) is of the same order as the corresponding data set. The vertical dashed and dotted lines represent the crossover times $T_2$ as in Eq. (10)

with

$$c = \frac{g - n}{(1 + n)(g - 1/2)}. \hspace{1cm} (19)$$

At the crossover, the infiltration length and its rate scale as

$$I_2 \sim r^{-c/2}$$

and the corresponding exponents $c/2$ are close to these predictions. In $d = 3$, Eq. (19) gives $c \approx 1.21$ ($c/2 \approx 0.61$). However, the same relation with $n_{eff}$ larger than the asymptotic $n$, as shown in Fig. 5(b), gives a smaller effective value of $c$. Indeed, the numerical estimates of exponent $c$ in the scaling of $T_2$ [Fig. 12(a)] are smaller than the estimate of the scaling approach and the numerical estimates of the exponents of $T_2$ and of $T_2$ [Figs. 11(b) and 11(c)] are smaller than the corresponding estimate of $c/2$, with differences in the range $9-13\%$.

The number of time decades $N_{tir}$ with time increasing rates is

$$N_{tir} \sim \log_{10} \left( \frac{T_2}{T_1} \right) \sim (c - b) \log_{10} r^{-1}, \hspace{1cm} (21)$$

where $c - b \approx 0.25$ in $d = 2$ and $c - b \approx 0.33$ in $d = 3$. In cases of very slow reactions or very fast diffusion, Eq. (3) implies $r \ll 1$ and the time increasing rates may be observed in several time decades.

For comparison, in the reactive regime of the dissolution of a compact medium, the constant rate lasts longer, with a number of time decades $\sim \log_{10} r^{-2}$. However, in that case, the crossover to Fickean infiltration occurs with continuously decreasing rates instead of the time increasing rates obtained here. The difference is related to the penetration of the reactants in the percolating porous system, which dissolve the pore walls and accelerate the convergence to the Fickean regime.

Our numerical results also show time increasing rates of the reaction lengths. However, we were not able to determine a scaling relation for that length. In Fig. 13 we show the effective slopes of the log $R \times \log T$ plots for three values of $r$ in $d = 3$. These slopes have peaks between $T_1$ and $T_2$, but they do not converge to a finite exponent as $r$ decreases; instead, they are increasing, which means that faster growths of $R$ and $R$ are observed in the regime of time increasing infiltration rate. Similar results are obtained in $d = 2$. Direct inspection of Figs. 7(b), 7(d), 10(b), and 10(d) also show that it is not possible to perform reliable linear fits of the log $R \times \log T$ or of the log $R \times \log T$ plots in that regime. Despite this limitation, we observe that $R_2 \sim I_2$ and $R_2 \sim I_2$ in both dimensions, which indicate that the reaction length crosses over to the Fickean regime simultaneously with the infiltration length.

**FIG. 13.** (Color online) Effective exponent of $R$ as a function of $T$. The vertical dashed and dotted lines represent the crossover times $T_1$ and $T_2$, respectively, with colors matching the corresponding data set.

**VII. CONCLUSION**

We studied a model of solute infiltration from a source of constant concentration into disordered lattices at the percolation thresholds and reactions of the solute with the impermeable sites, which create new porous sites. We performed numerical simulations in hypercubic lattices in $d = 2$ and 3 and developed scaling approaches to determine the time evolution of the extents of infiltration and reaction and to relate them with a model parameter $r$ that describes the relative rate of reaction and diffusion (Dankohler number). Cases of slow reactions ($r \ll 1$) are considered.

The model without reactions shows subdiffusive infiltration with a time scaling exponent $n$ predicted by
the same relation previously verified in infinitely ramified fractals. In the reactive case, short time subdiffusion is observed and, at sufficiently long times, the reacting media is far from the source and the infiltration is Fickian. Between these regimes, a regime with time increasing rates of infiltration and reaction is observed. This is explained by a cooperative effect between the directional infiltration of the solute into the fractal porous medium and the advance of reactions in all directions. The exponents of the time evolution of the infiltration and reaction lengths and their relations with $r$ are predicted by a scaling approach and confirmed by numerical simulations, with some deviations in $d = 3$ that can be explained by the slow convergence of the initial subdiffusion exponent. The regime with time increasing rates spans a time range that increases as $r$ decreases, so that it is more likely to be observed in slowly reacting materials.

The interplay of infiltration in porous media and chemical reactions that change their structures is essential to understand their evolution. This was already shown in the study of materials of geological and technological interest. In low porosity systems, diffusion is expected to be the main transport mechanism, and if those systems are fractal, they are expected to display subdiffusion. This work suggests the investigation of a possible anomalously fast infiltration if the reactions increase the porosity of such systems.

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