Exact calculation of the tortuosity in disordered linear pores in the Knudsen regime

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The squared reciprocal tortuosity $\kappa^{-2} = D/D_0$ for linear diffusion on lattices and in pores in the Knudsen regime is calculated analytically for a large variety of disordered systems. Here, $D_0$ and $D$ are the self diffusion coefficients of the smooth and the corresponding disordered system, respectively. To this end, a building block principle is developed that composes the systems into sub-structures without cross-correlations between them. It is shown how the solutions of the different building blocks can be combined to gain $D/D_0$ for pores of high complexity from the geometrical properties of the systems, i.e. from the volumes of the different sub-structures. As a test, numerical simulations are performed that agree perfectly with the theory.

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I. INTRODUCTION

Diffusion in random media has been a subject of large interest in the last decades (for some latest reports see e.g. [1]). In the last years, the interest has focused on the experimentally accessible subject of diffusion of gas molecules in pores (see Fig. 1), as e.g. the human lung [2], linear silicon nanochannels [3] or zeolites and other micro- and nanopores [4–6]. As recent progress in synthesizing nanostructured porous materials has provided the options of designing specific pore architectures [7], an exact analytical understanding of the diffusion process is of great importance. Of particular interest is the tortuosity factor $\kappa = \sqrt{D_0/D}$ that describes the relation between the diffusion coefficients $D$ and $D_0$ of systems with and without geometrical disorder [8]. Both, $D$ and $D_0$, can be gained by studying either the transport or the self-diffusion problem, where in the Knudsen regime, the self- and the transport diffusion coefficients are the same for a given geometry. Theoretical calculations of $D$ on complex pores have mostly been based on numerical simulations of the transport- [9–12] or the self diffusion problem [11–15] and/or phenomenological or semi-analytical approaches [16], whereas exact analytical results of specific pore geometries have only been provided along loopless curved one-dimensional paths [17] and for systems with dead ends [18, 19]. Whereas in loopless curved systems (see Fig. 2(b,e)), the tortuosity factor is determined by the longer path, the particle has to travel along the curve in order to overcome a smaller distance in $x$-direction, the diffusion time of a particle in pores with dead ends (as shown in Fig. 2(c,f)) is increased by detours into the dead ends that do not contribute to the diffusion along the $x$-direction.

The purpose of this work is to create a new approach, based on the self diffusional problem, for the exact analytical calculation of $D/D_0$ of more complex systems as the ones of Fig. 2.

FIG. 1: [Color online] Sketch of the diffusion process inside a (smooth) pore. The particle is reflected with different angles between the pore walls, leading to jump lengths of very different sizes.

FIG. 2: [Color online] Sketch of the geometries of special well-known sub-units for lattices (a–c), left column and the corresponding pores (d–f), right column, i.e. regular units (a,d), curved units (b,e) and systems with dead-end units (c,f). In the pores, all $x$-channels (red dashed lines) are of square cross section (with side length $h$) and up to 1000 of these (identical) blocks are sticked together to account for an infinite elongation into the $x$-direction. For further geometric details see caption of Tab. I.

To this end, we consider the lattice problem and its connection to diffusion in linear pores in the Knudsen regime [20] (see below). To calculate $D/D_0$, we decompose the considered complex systems (see Fig. 3 for examples) into simpler exactly solvable geometric sub-structures (building blocks) without cross-correlations between them and show how the results of the single sub-units must be combined to calculate $D/D_0$ as a function of simple geometrical data, i.e. of the different lengths, widths and volumes as given in Tab II. We verify our results by numerical simulations that agree perfectly with the theoretical predictions. In this work, we are only interested in systems where a fully analytical treatment is possible, i.e. where all individual building blocks can be solved analytically. However, we would like to point out that this method can also go beyond these cases by combining analytical and numerical data of different building blocks.

FIG. 3: [Color online] Example of a building block for a pore in the Knudsen regime of complicated geometry. This system is built up of $N$ (a) part of a smooth linear pore, (b) part of a linear dead end, (c) two linear dead ends connected, (d) a linear dead end connected to the large linear one, (e) part of a loopless curved horizontal path, (f) part of a loopless curved diagonal path.
The paper is organized as follows: In section II, we present the random walk on a lattice and the diffusion problem in pores, while the underlying theory for the calculation of $D$ for systems made of various building blocks is explained in section III. In section IV, we present the theoretical results and verify them by numerical simulations. In the last section V, we discuss the results and give an outlook.

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<th>Fig.</th>
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TABLE I: Table of the geometries and the analytical results of the simple units from Fig. 2. Upper half: lattice systems, lower half: pores. All units are the same as in Tab. II.

II. DIFFUSION ON LATTICES AND IN PORES

In a linear random walk, a particle jumps inside a $d$-dimensional lattice (see left column of Figs. 2 and 3) and we are interested in its displacement in $x$-direction. We concentrate on problems, where despite an irregular structure of the systems the long-time diffusion stays normal, which means that $D$ for long times is defined by the Einstein relation,

$$\lim_{t\to\infty} \langle x^2(t) \rangle = 2Dt,$$

where the mean square displacement $\langle x^2(t) \rangle$ is the squared distance, a particle has traveled during time $t$ in $x$-direction. For anomalous diffusion, as e.g. on fractal structures, we refer to the literature [21–24]. For simplicity, we concentrate on cubic (square) lattices, where in the absence of disorder each lattice site has $2d$ neighbors and on unbiased walks, where jumps to the neighboring sites occur with equal probability. Disorder is created by the removal of sites or of links between neighboring sites. On a lattice, a walker chooses one of the $2d$ possible directions for the following jump at random. If the link to the chosen neighbor is existing, the walker jumps, thereby performing a jump of length $a$ (lattice constant) during a time step $\tau$. If the link has been removed, the walker stays for this time step where it is (waiting time).

Diffusion in pores (right column of Figs. 3 and 2) represents a more complex problem where, in general, the track of the gas molecules through the pores depends on the collisions between the gas molecules as well as on the collisions of the gas with the pore walls. In cases where Knudsen diffusion [20] dominates, as it has been shown in various transport

FIG. 3: [Color online] Geometries for lattices (left column) and the corresponding pores (right column) that are analyzed by the building block principle: (a, e) curved system with dead ends, (b, f) curved system in parallel with a channel, (c, g) system of two intersecting curves. All together to account for an infinite elongation into the $x$-direction for further geometric details see Tab. II.)

TABLE II: Table of the geometries and analytical results based of the combined systems of several building blocks from Fig. 3. Upper half: lattice systems, lower half: pores. All lengths on lattices and in pores are given in units of the lattice constant $a$ and the pore diameter $h$, respectively. $D$ is referred to the value $D_0 = a^2/(4\tau)$ for the lattice ($d = 2$) and to $D_0 = 0.37 \hbar v_0$ for the pores. All omitted numbers are equal to 1.

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TABLE II: Table of the geometries and analytical results based of the combined systems of several building blocks from Fig. 3. Upper half: lattice systems, lower half: pores. All lengths on lattices and in pores are given in units of the lattice constant $a$ and the pore diameter $h$, respectively. $D$ is referred to the value $D_0 = a^2/(4\tau)$ for the lattice ($d = 2$) and to $D_0 = 0.37 \hbar v_0$ for the pores. All omitted numbers are equal to 1.
situations through porous media [3, 25], the interactions of the molecules with the pore walls play the crucial role and the intermolecular collisions can be neglected. In this case, the molecules perform a series of free flights and change the flight direction independently from each other after collisions with the pore walls as shown in Fig. 1. Therefore the problem is reduced to many independent individual flights. In this work, we concentrate on Knudsen diffusion under Lambert’s reflection law in three-dimensional regular and irregular pores [4, 11, 12, 15]. In this picture, the particle is absorbed from the wall after collision and after a very short time (that is neglected) re-emitted into a random direction, where the new direction \( \theta \in [-\pi/2, \pi/2] \) to the normal component of the surface occurs with probability \( d\Omega = \cos \theta \, d\Omega \), where \( d\Omega = \sin \theta d\varphi d\varphi \) in \( d = 3 \).

It is clear that disorder slows down the diffusion process, leading to a smaller value of \( D \) as compared to \( D_0 \) of a smooth system. Quantitative calculations that connect \( D/D_0 \) to simple geometrical properties, as volumes and lengths of the different segments exist for loopless curved geometries and for systems with dead ends (dangling bonds) that are connected to the main channel by a thin entry, examples of which are both shown in Fig. 2.

In loopless curved geometries [17] (see Fig. 2(b,e)), the effective length \( \ell \) (also called “chemical length” [23, 24]) of the path a particle has to travel in order to come from \( A \) to \( B \) is larger than the \( x \)-distance between the same points (see Fig. 4 for an illustration). Therefore, normal diffusion with \( \langle x^2(t) \rangle = D_0 t \) applies for the effective length and with the relation \( \langle x(t) \rangle = \langle x(t') \rangle \) between \( x \)- and \( t \)-space, one finds \( D = D_0 \langle x(t) \rangle / \langle x(t) \rangle = D_0 (V_x/V)^2 \) [17], where the last expressions refers to pores with \( V_x \) and \( V \) as defined below.

In dead-end geometries (see Fig. 2(c,f)), as it was first discussed in [26], the walker only proceeds in the \( x \)-channels (indicated by the red dashed lines in Fig. 2), while the time inside the dead ends increases the total time \( t \) of the walk without increasing \( \langle x^2 \rangle \). Quantitative considerations [18, 19, 22] show that \( D/D_0 = V_x/V \), with the volume \( V_x \) of the \( x \)-channels and the system volume \( V \) (of channel plus dead ends).

In this work, we want to combine these well-known systems to more complex geometries by connecting them using additional segments or by intersecting them directly with each other, thereby forming networks. To this end, we show how a system of different sub-units, where the diffusion may be (i) uncorrelated (as in a straight channel), (ii) strongly correlated (as in dead ends, where each jump is compensated by a jump into the opposite direction) and (iii) intermediate correlated (as in curved channels, where correlated forward-backward jumps occur at all times) must be combined to obtain \( D/D_0 \) and thus \( D \). The approach uses the self diffusion picture but clearly, as the self- and transport diffusion coefficients of a given geometry are equal, is also valid for transport diffusion.

III. CALCULATION OF THE DIFFUSION COEFFICIENT

A. General Considerations

Generally, we can write \( \langle x^2(t) \rangle \) after \( N \) time steps as

\[
\langle x^2(t) \rangle = \left( \sum_{i=1}^{N} \xi_i^2 \right) = \left( \sum_{i=1}^{N} \xi_i^2 \right) + 2 \sum_{i,j=1}^{N} \xi_i \xi_j,
\]

(2)

where \( \xi_i \) and \( \xi_j \) are the single jump lengths in \( x \)-direction. In the following, we call the 1st term of the rhs of Eq. (2) the “quadratic term” and the 2nd term the “correlation term”.

To calculate \( D \) directly from the geometry of the system, we refer to the well-known principles that (i) the particle concentration (as well as the gas pressure) is identical all over the system and (ii) correlations among different walks do not exist (Knudsen condition). Condition (i) tells us that in the average over many walks, all places of the system are visited with equal probability. This is true for real experiments as well as for computer simulations, provided that the starting point is chosen with equal probability among all sites.

As the sequential order of the single time steps does not play a role for evaluating the quadratic term of Eq. (2), the single steps of the sum, even if they belong to different walks may be interchanged. Then, we can replace the time average of the quadratic term by the ensemble average and describe it solely by all jumps that occur at the same time on all places, i.e., by the geometric properties of the system and independently of the track of the single walks. We thus replace the quadratic term by \( \langle \sum_{i=1}^{N} \xi_i^2 \rangle = N \langle \xi^2 \rangle \), where \( \langle \xi^2 \rangle \) is the mean quadratic jump length in \( x \)-direction over all \( N \) jumps. The total time of the walk is \( t = N \langle t \rangle \), with the average duration of the time steps \( \langle t \rangle \). Jumps into the \( y \)- and \( z \)-direction count as waiting times, as they increase \( t \) without increasing \( x \).

On lattices, all time steps are equal and \( \langle t \rangle = \tau \), whereas \( \langle \xi^2 \rangle \) depends on the number of waiting times. For the diffusion coefficient \( D_0 \) of a \( d \)-dimensional ordered lattice of lattice constant \( a \) (where the correlation term is zero), we find \( \langle \xi^2 \rangle = a^2/d \) and therefore \( D_0 = a^2/(2d\tau) \). On a disordered lattice, on the other hand, some jump-trials into the \( x \)-direction find no bond and lead to additional waiting times. Furthermore, the correlation terms may give an additional negative contribution and accordingly, \( D < D_0 \).

In pores, the jump lengths and time steps are not constant and therefore \( D_0 \) depends on the cross-section of the pore. For the smooth pore with square surface section of side length \( h \) (see Fig. 2(d)), \( D_0 = \langle \xi^2 \rangle / \langle t \rangle \) is numerically found as \( \approx 0.37 h/v_0 \), with the velocity \( v_0 \) of the particle along the trajectory (see also [12]) and with \( h \) as unit length. (The time steps can be defined as \( \tau = h/v_0 \).) The generalization to circular or rectangular cross sections is straightforward, but not the
purpose of the present work. Therefore, we refer all values of $D$ in pores to the value $D_0$ of the corresponding smooth pore with unit length $h$ of Fig. 2(d) that we use as the basic element. Smooth pores with side length $nkh$ possess the diffusion coefficient $n_kD_0$.

B. The building block principle

We now turn to more complex geometries. The first step for a rigorous treatment is the decomposition of the considered system into well-known analytically treatable sub-units $k$ as e.g. $A$: smooth $x$-channels, $B$: loopless curved units, $C$: dead ends and $D$: vertical links. The sub-units must be chosen without cross-correlations, i.e. such that the steps a walker performs in a given sub-unit do not influence the steps in another unit or during a second visit of the same unit.

The relative time $t_k/t$, the particles spend in a given unit $k$ can be expressed by $t_k/t = N_k/N = V_k/V$, with the relative number of sites $N_k/N$ (in lattices) or the relative volume $V_k/V$ (in pores) of the $k$th unit. We also need the relative time, the particles spend in the $x$-channels (indicated by the red dashed lines in Figs. 2 and 3), $t_{x,k}/t = N_{x,k}/N = V_{x,k}/V$, where $N_{x,k}$ and $V_{x,k}$ are the number of sites and the volume of the $x$-channels of the $k$th unit, respectively. Dead-ends, even if oriented in $x$-direction do not count as $x$-channels. In the absence of cross-correlations between the different units, $(x(t)^2)$ can be gained from Eq. (2) by sum over all quadratic terms and correlation terms of all units $k$. As the mean quadratic jump length $\langle \xi^2 \rangle_2$ of the $k$th unit normally grows quadratically with the pore thickness $n_kh$ (except for possible cut-off effects in the jump-length distribution, see below) we have $\langle \xi^2 \rangle_k = n_k^2 \langle \xi^2 \rangle$ and get for the quadratic terms with $t = N(t)$

$$
\sum_i N_i \langle \xi^2 \rangle_i/N = \sum_k n_kD_0N_{x,k}/N = \sum_k n_kD_0V_{x,k}/V,
$$

where the index $i$ runs over the time steps and $k$ runs over the geometric sub-units. For simplicity, we investigate networks with units of identical thickness here, i.e. all $n_k = 1$. If all correlation terms of all units are equal to zero, as e.g. for a system of parallel pores, $D/D_0$ can be gained from (3). Otherwise, we also have to calculate the correlation terms of Eq. (2), which can be done analytically for several types of building blocks. Here, we treat the following cases:

(A): The simplest units are regular ordered lattices and straight channels (see Fig. 2(a,d)), where each jump is followed by a positive or a negative $x$-jump with equal probability. It is common knowledge that the correlation terms of these sub-units are 0.

(B): "Curved geometries" (see Fig. 2(b,e)) consist of one loopless curved backbone, where the particles perform a large number of forward-backward jumps before they pass the corners. So, the $x$-jumps are correlated, because positive $x$-jumps to corner sites are more likely followed by negative (than by other positive) $x$-jumps. As described above, $D$ can be calculated by considering the problem in $t$-space, where $t_k$ is the effective or chemical length of the curve $k$. So, if the $k$th unit

is a loopless curve, $D$ of this unit alone can be written as [17]

$$
D = D_0 \left( \frac{x_k}{t_k} \right)^2 = D_0 \left( \frac{V_{x,k}}{V_k} \right)^2,
$$

where for the case of pores, $t_k$ and $x_k$ have been replaced by the total volume $V_k$ and the $x$-volume $V_{x,k}$. From Eqs. (1)-(4) we find the correlation term of the loopless curves,

$$
2 \sum_{i,j,k}^N \xi_i \xi_j = 2t_k \left( D - D_0 \frac{V_{x,k}}{V_k} \right) = 2t_kD_0 \left( \frac{V_{x,k}}{V_k} \right)^2 - 2t_kD_0 \frac{V_{x,k}}{V_k},
$$

(C): The simplest correlated sub-units are dead ends [19, 26], i.e. units that are connected to the other parts by a thin entry (see Fig. 2(c,f)). The entry and the exit point to the dead ends coincide or otherwise speaking, the whole path inside the dead ends is considered as pure delay. This means that the correlation term cancels with the quadratic term, so that here, the diffusion is suppressed and both terms can be set to zero (even if, strictly speaking, none of them is zero, but one term has the negative value of the other).

(D): In straight $z$-or $y$-paths ("vertical links"), $x$-jumps are not possible and therefore the quadratic as well as the correlation term are both equal to zero. Vertical links only increase $t$ without increasing $x$ and therefore act in the same way as dead ends. Indeed, these paths need not even be completely straight – it is sufficient if the lower and the upper entrance point possess the same $x$-value.

We now turn to complex systems that can be considered as combinations of the described systems. We discuss in which way the sub-units (A)-(D) can be combined without generating cross-correlations between them, so that the combined geometries can be simply treated by summing over all quadratic and all correlation terms of all blocks.

(i) Dead ends (C) can be added to all systems at arbitrary positions (see Fig. 3(a,e)). As we have seen, they don’t bring new terms, but increase $t$. (ii) Several infinite units, e.g. (A) and (B), can be connected by vertical links (D) to form a simple network of interconnected parallel pores (see Fig. 3(b,f)). There are no cross-correlations between (A) and (B) as long as both are infinitely elongated into the $x$-direction: if a random walker changes from (A) to (B) (or vice versa) it can continue its path in (B) into both directions with equal probability, so that no step in (A) influences later steps in (B) and vice-versa. Also $x$-correlations between (A) and (D) (or (B) and (D)) cannot exist, as $x$-jumps in (D) either do not exist (if the link is completely straight) or exist in pairs where positive and negative steps always cancel. Therefore terms $(x,t,x)$ with $x_i$ or $x_j$ in (C) are either zero or appear with a negative counterpart. So, also systems of several units (A) and/or (B) connected by vertical links can be calculated by the building-block principle. (iii) A more complex combination appears, when two infinite units (A) and (B) are intersected without
intermediate pieces as shown in Fig. 3(c,d,g,h). Again, no cross-correlations between (A) and (B) take place (by the same argument as before) and the total system can again be treated by the building-block principle. (iv) Unfortunately not treatable this way are structures as shown in Fig. 5(a) involving finite pieces creating short-cuts inside the same structure (B), because in those cases new correlations are created (inside (B) as well as between the finite units). (v) Also not-treatable in this way are systems where the connection between two sub-units contains finite x-paths, i.e., where the lower and the upper entry point do not have the same x-value (see Fig. 5(b)), because in those cases cross-correlations between the different finite units take place. We would like to point out, however, that in both cases new sub-units could be defined, calculated numerically and then combined with the analytical results of other exactly treatable building blocks. But this goes beyond the purpose of the present paper.

In summary, there is a large variety of combinations of well-known building blocks, leading to quite complex geometries, from which \( \frac{D}{D_0} \) can be calculated analytically which we will show in detail in section IV and verify by numerical simulations. The method is equally applicable to random walks on lattices and to diffusion in pores.

C. Numerical calculations

The analytical calculations (explained in the next section) are compared to numerical simulations on the systems shown in Figs. 2 and 3. The particle flow takes place along the x-direction and the figures of the geometries are meant to be infinitely elongated along the x-axis. The simulations have been performed for different geometric details (as listed in the tables) and 100 – 1000 elementary units have been stucked together. For technical reasons, namely for a faster generation of the systems in the computer simulations and for the presentation of the results in tables, the systems are periodic, but all calculations are valid for completely disordered structures as well. In order to choose the starting point among all lattice sites with equal probability, periodic boundaries should be chosen in non-periodic systems to enable walks of arbitrary length to both sides of the starting point. In the periodic systems calculated here, it is sufficient to choose the starting point with equal probability among all sites of one unit.

The computation of the random walk on a lattice is well-known, so that we refer to the literature (see e.g. \[23, 24\]). For the diffusion processes through pores, all walls of a given geometry have been stored and ordered according to their position and orientation (normal vector \( \vec{n} \)). For the computations of the particle flights from wall to wall with a given direction, the possible intersections of the flight trajectory with walls of increasing distance from the particle position have been computed. Once an intersection point is found, the computation can be stopped for all walls of the same \( \vec{n} \) but with larger distance to the actual particle position. The computation of \( \langle x^2(t) \rangle \), once the collision point with the next wall has been found, is straightforward and takes place as on a lattice.

Note that for the sub-unit of the curved systems (Fig. 2(e)), some specialities exist in pores (as compared to lattices) that we mention briefly: First, due to Lambert’s reflexion law, the probability for long jumps into the x-direction is slightly different when starting on a vertical and on a horizontal wall, which should influence the results of rough systems slightly. However, in three-dimensional pores, these differences are not large, so that we neglect them here. They would be more important in \( 2d \)- pores, see Ref. [11]). Second, finite x-segments lead to an upper cut-off of the jump lengths distribution \( P(\xi) \) and therefore to a modified value of \( \langle x^2(t) \rangle \) that cannot be taken into account analytically. Clearly, the correct jump length distribution could be determined numerically for each system, but this is not the aim of the present work. Therefore, we choose systems with lengths of the x-segments larger than the average jump length of the smooth system, so that both distributions \( P(\xi) \) are nearly the same. The other segments (oriented along the y- or z-direction) may be of arbitrary lengths because there, the cut-off of the jump-lengths leads to additional factors in the relation \( \langle x^2(t) \rangle = D_{\delta t} \) and Eq. (4) that cancel.

We test the numerical calculations on the simple units of Fig. 2, i.e. on one smooth system, one system with dead ends and one curved system and compare them to Eqs. (3) and (4), respectively. The results are shown in Fig. 6 for lattices and pores and agree perfectly with the expectations.

IV. NETWORKS OF DIFFERENT BUILDING BLOCKS

In this section, we apply the building block principle to the geometries of Fig. 3. To this end, we need for each sub-unit \( k \) the number of sites \( N_k \) or the volume \( V_k \) (for lattices and pores, respectively) as well as the number \( N_{\alpha,k} \) or the volume \( V_{\alpha,k} \) of...
As the last example, we intersect two curves with volumes \( V_1 \equiv V_{C1}, V_2 \equiv V_{C2} \), \( x \)-volumes \( V_{x1} \equiv V_{x,C1}, V_{x2} \equiv V_{x,C2} \); and \( V = V_{C} + V_{C2}, V_{x} = V_{x,C1} + V_{x,C2} \). We get \( D/D_0 \) by adding the quadratic and the correlation terms of both systems, yielding

\[
D = D_0 \frac{V_{x,C}^2}{VV_C} + D_0 \frac{V_{x,C}^2}{VV_C}.
\]  

We can see that in all described cases, the ratio \( D/D_0 \) can be directly obtained from purely geometrical data without performing numerical simulations. If we consider lattices instead of pores, all values of \( V, V_1, V_2, V_{x,1}, \) and \( V_{x,2} \) have to be replaced by the respective values of \( N, N_1, \) and \( N_{x,1,2} \).

Nevertheless, we performed numerical simulations over an average of \( 10^5 \) systems (except for the pore systems of Eq. (7) and (8), where due to larger calculation times an average over only \( 10^3 \) systems has been performed) to put the relations (6)-(8) to a direct test. The results of the simulations (symbols) are shown in Fig. 7 and compared to the theoretical values (straight lines) for systems of different geometrical details as listed in Tab. II. The figures show the results for lattices (Fig. 7(a)) as well as for pores (Fig. 7(b)) and in all considered cases, the agreement between numerical and theoretical data is excellent. (Larger fluctuations in the pore realizations of the two last systems are due to the poorer statistics.) Clearly, also more than two units can be combined and additional dead ends can be easily included to all considered systems to increase \( V \), i.e. the same calculation scheme can also be applied to various other geometries, including real large networks.

V. CONCLUSION AND OUTLOOK

We have presented an analytical method to calculate the tortuosity factor \( \kappa \) that describes the decrease of the diffusion coefficient \( D \) in the presence of disorder as compared to the diffusion coefficient \( D_0 \) of a smooth system for a large variety of complex disordered systems. To this end, we have developed a building block principle that is based on a careful analysis of the correlation effects of the diffusion process. For the systems of this work, we could express \( \kappa \) simply by the volumes of the different sub-structures. The procedure has been demonstrated on many different systems and it is clear that many more systems can be constructed accordingly.

We only considered systems, where an analytical treatment is possible for both, the random walk on lattices and for the pore diffusion. However, the building block principle can also be combined with correlations obtained from numerical simulations, as e.g. dangling bonds with a thickness \( \delta > h \) (where the correlation term does not exactly cancel with the quadratic term), curved systems where the \( x \)-paths are small or backbones of varying thickness. So, additionally to the systems considered here, a large variety of pores can be manufactured and understood by combining building blocks of numerically and exactly obtained correlation terms along the lines of the present paper.

We also pointed out that not all types of combinations between sub-units are suited for this treatment – if cross-correlations between the sub-units are created, the described
method fails. We showed several examples, where this is the case. However, a large variety of quite complex systems can be treated in the way described and it is straightforward to cross e.g. many loopless curved systems to get real networks that can be calculated by this building block principle.

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