Phase Transitions on Fractals and Networks

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Glossary

Cluster
Clusters are sets of occupied neighbouring sites.

Critical exponent
At a critical point or second-order phase transition, many quantities diverge or vanish with a power law of the distance from this critical point; the critical exponent is the exponent for this power law.

Diffusion
A random walker decides at each time step randomly in which direction to proceed. The resulting mean square distance normally is linear in time.

Fractals
Fractals have a mass varying with some power of their linear dimension. The exponent of this power law is called the fractal dimension and is smaller than the dimension of the space.

Ising model
Each site carries a magnetic dipole which points up or down; neighbouring dipoles “want” to be parallel.
Percolation
Each site of a large lattice is randomly occupied or empty.

1 Definition and Introduction

Some phase transitions, like the ferromagnetic Curie point where the spontaneous magnetisation vanishes, happen in solids, and experiments often try to grow crystals very carefully such that the solid in which the transition will be observed is periodic with very few lattice faults. Other phase transitions like the boiling of water or the liquid-vapour critical point, where the density difference between a liquid and its vapour vanishes, happen in a continuum without any underlying lattice structure. Nevertheless, the critical exponents of the Ising model on a simple-cubic lattice agree well with those of liquid-vapour experiments. Impurities, which are either fixed (“quenched dilution”) or mobile (“annealed dilution”), are known to change these exponents somewhat, e.g. by a factor $1 - \alpha$, if the specific heat diverges in the undiluted case at the critical point, i.e. if the specific heat exponent $\alpha$ is positive. In this review we deal neither with regular lattices nor with continuous geometry, but with phase transitions on fractal and other networks. We will compare these results with the corresponding phase transitions on infinite periodic lattices like the Ising model.

2 Ising Model

Ernst Ising in 1925 (then pronounced EEsing, not EYEsing) published a model which is, besides percolation, one of the simplest models for phase transitions. Each site $i$ is occupied by a variable $S_i = \pm 1$ which physicists often call a spin but which may also be interpreted as a trading activity \[1\] on stock markets, as a “race” or other ethnic group in the formation of city ghettos \[2\], as the type of molecule in binary fluid mixtures like isobutyric acid and water, as occupied or empty in a lattice-gas model of liquid-vapour critical points, as an opinion for or against the government \[3\] \[4\], or whatever binary choice you have in mind. Also models with more than two choices, like $S_i = -1, 0$ and $1$ have been investigated both for atomic spins as well as for races, opinions, .... Two spins $i$ and $k$ interact with each other by an energy $-JS_iS_k$ which is $-J$ if both spins are the same and $+J$ if they
are the opposite of each other. Thus $2J$ is the energy to break one bond, i.e. to transform a pair of equal spins to a pair of opposite spins. The total interaction energy is thus

$$E = -J \sum_{<i,k>} S_i S_k,$$

(1a)

with a sum over all neighbour pairs. If you want to impress your audience, you call this energy a Hamiltonian or Hamilton operator, even though most Ising model publications ignore the difficulties of quantum mechanics except for assuming the discrete nature of the $S_i$. (If instead of these discrete one-dimensional values you want to look at vectors rotating in two- or three-dimensional space, you should investigate the XY or Heisenberg models instead of the Ising model.)

Different configurations in thermal equilibrium at absolute temperature $T$ appear with a Boltzmann probability proportional to $\exp(-E/k_B T)$, and the Metropolis algorithm of 1953 for Monte Carlo computer simulations flips a spin with probability $\exp(-\Delta E/k_B T)$, where $k_B$ is Boltzmann’s constant and $\Delta E = E_{\text{after}} - E_{\text{before}}$ the energy difference caused by this flip. If one starts with a random distribution of half the spins up and half down, using this algorithm at positive but low temperatures, one sees growing domains. Within each domain, most of the spins are parallel, and thus a computer printout shows large black domains coexisting with large white domains. Finally, one domain covers the whole lattice, and the other spin orientation is restricted to small clusters or isolated single spins within that domain. This self-organisation (biologist may call it “emergence”) of domains and of phase separation appears only for $0 < T < T_c$ and only in more than one dimension. For $T > T_c$ (or at all positive temperatures in one dimension) we see only finite domains which no longer engulf the whole lattice. This phase transition between long-range order below and short-range order above $T_c$ is called the Curie or critical point; we have $J/k_B T_c = \frac{1}{2} \ln(1 + \sqrt{2})$ on the square lattice and 0.221655 on the simple cubic lattice with interactions to the $z$ nearest lattice neighbours; $z = 4$ and 6, respectively. The mean field approximation becomes valid for large $z$ and gives $J/k_B T_c = 1/z$. Near $T = T_c$ the difference between the number of up and down spins vanishes as $(T_c - T)^\beta$ with $\beta = 1/8$ in two, $\simeq 0.32$ in three, and $1/2$ in six and more dimensions and in mean field approximation.
We may also influence the Ising spins though an external field $h$ by adding

$$-h \sum_i S_i$$

(1b)

to the energy of Eq.(1a). This external field then pushes the spins to become parallel to $h$. Thus we no longer have emergence of order from the interactions between the spins, but imposition of order by the external field. In this simple version of the Ising model there is no sharp phase transition in the presence of this field; instead the spontaneous magnetisation (fraction of up spins minus fraction of down spins) smoothly sinks from one to zero if the temperature rises from zero to infinity.

3 Fractals

Fractals obey a power law relating their mass $M$ to their radius $R$:

$$M \propto R^D$$

(2)

where $D$ is the fractal dimension. An exactly solved example are random walks (= polymer chains without interaction) where $D = 2$ if the length of the walk is identified with the mass $M$. For self-avoiding walks (= polymer chains with excluded volume interaction), the Flory approximation gives $D = (d + 2)/3$ in $d \leq 4$ dimensions ($D(d \geq 4) = 2$ as for random walks), which is exact in one, two and four dimensions, and too small by only about two percent in three dimensions.

We now discuss the fractal dimension of the Ising model. In an infinite system at temperatures $T$ close to $T_c$, the difference $M$ between the number of up and down spins varies as $(T_c - T)^\beta$ while the correlation length $\xi$ varies as $|T - T_c|^{-\nu}$. Thus, $M \propto \xi^{-\beta/\nu}$. The proportionality factor varies as the system size $L^d$ in $d$ dimensions since all spins are equivalent. In a finite system right at the critical temperature $T_c$ we replace $\xi$ by $L$ and thus have $M \propto L^{d-\beta/\nu} = L^D$ with the fractal dimension

$$D = d - \beta/\nu \quad (d \leq 4)$$

(3a)

Warning: one should not apply these concepts to spin clusters if clusters are simply defined as sets of neighbouring parallel spins; to be fractals at $T = T_c$ the clusters have to be sets of neighbouring parallel spins connected
by active bonds, where bonds are active with probability $1 - \exp(-2J/k_BT)$. Then the largest cluster at $T = T_c$ is a fractal with this above fractal dimension.

This warning is no longer valid for percolation theory (see separate reviews in this encyclopedia) where each lattice site is occupied randomly with probability $p$ and clusters are defined as sets of neighbouring occupied sites. For $p > p_c$ one has an infinite cluster spanning from one side of the sample to the other; for $p < p_c$ one has no such spanning cluster; for $p = p_c$ one has sometimes such spanning clusters, and then the largest or spanning cluster has

$$M \propto L^D; \quad D = d - \beta \nu \quad (d \leq 6)$$

with the critical exponents $\beta$, $\nu$ of percolation instead of Ising models.

These were probabilistic fractal examples, as opposed to deterministic ones like the Sierpinski carpets and gaskets, which approximate in their fractal dimensions the percolation problem. We will return to them in the section “Ising models on fractals”.

Now, instead of asking how phase transitions produce fractals we ask what phase transitions can be observed on these fractals.

\section{4 Diffusion on Fractals}

\subsection{4.1 Unbiased diffusion}

The most thoroughly investigated phase transitions on fractals are presumably random walkers on percolation clusters \[31\], particularly at $p = p_c$. This research was started by Brandt \[8\] but it was the later Nobel laureate de Gennes \[9\] who gave it the catchy name “ant in the labyrinth”. The anomalous diffusion \[5, 6, 7\] then made it famous a few years later and may have also biological applications \[10\].

We put an ant onto a randomly selected occupied site in the middle of a large lattice, where each site is permanently occupied (randomly with probability $p$) or empty $(1 - p)$. At each time step, the ant selects randomly a neighbour direction and moves one lattice unit in this direction if and only if that neighbour site is occupied. We measure the mean distance

$$R(t) = <r(t)^2>^{1/2} \quad \text{or} \quad = <r(t)>$$
Figure 1: Log-log plot for unbiased diffusion at (middle curve), above (upper data) and below (lower data) the percolation threshold $p_c$. We see the phase transition from limited growth at $p_c - 0.01$ to diffusion at $p_c + 0.01$ separated by anomalous diffusion at $p_c$. Average over 80 lattices with 10 walks each.

where $\mathbf{r}$ is the vector from the starting point of the walk and the present position, and $r = |\mathbf{r}|$ its length. The average $< \ldots >$ goes over many such walking ants and disordered lattices. These ants are blind, that means they do not see from their old place whether or not the selected neighbour site is accessible (occupied) or prohibited (empty). (Also myopic ants were grown which select randomly always an occupied neighbour since they can see over a distance of one lattice unit.) The squared distance $r^2$ is measured by counting how often the ant moved to the left, to the right, to the top, to the bottom, to the front, or to the back on a simple cubic lattice.

The problem is simple enough to be given to students as a programming project. They then should find out by their simulations that for $p < p_c$ the above $R$ remains finite while for $p > p_c$ is goes to infinity as $\sqrt{t}$, for sufficiently long times $t$. But even for $p > p_c$ it may happen that for a single
ant the distance remains finite: If the starting point happened to fall on a finite cluster, then $R(t \to \infty)$ measures the radius of that cluster. Right at $p = p_c$, instead of a constant or a square-root law, we have anomalous diffusion:

$$R \propto t^k, \quad k = (\nu - \beta/2)/(2\nu + \mu - \beta)$$

(5)

for sufficiently long times. This exponent $k$ is close to but not exactly $1/3$ in two and $1/5$ in three dimensions. $\beta$ and $\nu$ are the already mentioned percolation exponents, and $\mu$ is the exponent for the conductivity if percolation is interpreted as a mixture of electrically conducting and insulating sites. If we always start the ant walk on the largest cluster at $p = p_c$ instead of on any cluster, the formula for the exponent $k$ simplifies to $\nu/(2\nu + \mu - \beta)$. The theory is explained in detail in the standard books and reviews [30, 31]. We see here how the percolative phase transition influences the random walk and introduces there a transition between diffusion for $p > p_c$ and finite motion.
Figure 3: Difficulties at transition from drift (small bias, upper data) to slower motion (large bias, lower data); 80 lattices with 10 walks each.

for $p < p_c$, with the intermediate “anomalous” diffusion (exponent below 1/2) at $p = p_c$. Fig.1 shows this transition on a large cubic lattice.

4.2 Biased diffusion

Another type of transition is seen in biased diffusion, also for $p > p_c$. Instead of selecting all neighbours randomly, we do that only with probability $1 - B$, while with probability $B$ the ant tries to move in the positive $x$-direction. One may think of an electron moving through a disordered lattice in an external electric field. For a long time experts discussed whether for $p > p_c$ one has a drift behaviour (distance proportional to time) for small $B$, and a slower motion for larger $B$, with a sharp transition at some $p$-dependent $B_c$. In the drift regime one may see log-periodic oscillations $\propto \sin(\text{const} \log t)$ in the approach towards the long-time limit, Fig.2. Such oscillations have been predicted for stock markets [11], where they could have made us rich, but
for diffusion they hamper the analysis. They come presumably from sections of occupied sites which allow motion in the biased direction and then end in prohibited sites [12].

Even in a region without such oscillations, Fig. 3 shows no clear transition from drift to no drift; that transition could only be seen by a more sophisticated analysis which showed for the $p$ of Fig. 3 that the reciprocal velocity, plotted versus log(time), switches from concave to convex shape at $B_c \simeq 0.53$. Fortunately, only a few years after these simulations [13] the transition was shown to exist mathematically [14].

These simulations were made for $p > p_c$; at $p = p_c$ with a fractal largest cluster, drift seems impossible, and for a fixed $B$ the distance varies logarithmically, with a stronger increase slightly above $p_c$ and a limited distance slightly below $p_c$, Fig. 4.
5 Ising Model on Fractals

What happens if we set Ising spins onto the sites of a fractal? In particular, but also more generally, what happens to Ising spins on the occupied site of a percolation lattice, when each site is randomly occupied with probability \( p \)? In this case one expects three sets of critical exponents describing how the various quantities diverge or vanish at the Curie temperature \( T_c(p) \). For \( p = 1 \) one has the standard Ising model with the standard exponents. If \( p_c \) is the percolation threshold where an infinite cluster of occupied sites starts to exist, then one has a second set of exponents for \( p_c < p < 1 \), where \( 0 < T_c(p) < T_c(p = 1) \). Finally, for zero temperature as a function of \( p - p_c \) one has the percolation exponents as a third set of critical exponents. (If \( p = p_c \) and the temperature approaches \( T_c(p_c) = 0 \) from above, then instead of powers of \( T_c(p_c) \) exponential behaviour is expected.) In computer simulations, the second set of critical exponents is difficult to observe; due to limited accuracy the effective exponents have a tendency to vary continuously with \( p \).

The behaviour at zero temperature is in principle trivial: each cluster of occupied neighbor has parallel spins, the spontaneous magnetisation is given by the largest cluster while the many finite cluster cancel each other in their magnetisation. However, the existence of several infinite clusters at \( p = p_c \) disturbs this argument there; presumably the total magnetisation (i.e. not normalised at magnetisation per spin) is a fractal with the fractal dimension of percolation theory.

Deterministic fractals, instead of the random “incipient infinite cluster” at the percolation threshold, may have a positive \( T_c \) and then allow a more usual study of critical exponents at that phase transition. Koch curves and Sierpinski structures have been intensely studied in that aspect since decades. To build a Sierpinski carpet we take a square, divide each side into three thirds such that the whole square is divided into nine smaller squares, and then we take away the central square. On each of the remaining eight smaller squares this procedure is repeated, diving each into nine squarelets and omitting the central squarelet. This procedure is repeated again and again, mathematically ad infinitum. Physicists like more to think in atoms of a fixed distance and would rather imagine each square to be enlarged in each direction by a factor three with the central square omitted; and then again and again this enlargement is repeated. In this way we grow a large structure built by squares of unit area.
Unfortunately, the phase transitions on these fractals depend on details and are not already fixed if the fractal dimension is fixed. Also other properties of the fractals like their “ramification” are important [15]; see [16] for recent work. This is highly regrettable since modern statistical physics is not restricted to three dimensions. Models were studied also in seven and in minus two dimensions, in the limits of dimensionality going to infinity or to zero, or for non-integral dimensionality. (Similarly, numbers were generalized from positive counts to negative integers, to rational and irrational numbers, and finally to imaginary/complex numbers.) It would have been nice if these fractals would been models for these non-integral dimensions, giving one and the same set of critical exponents once their fractal dimension is known. Regrettably, we had to give up that hope.

Many other phase transitions, like those of Potts or voter models, were studied on such deterministic fractals, but are not reviewed here. We mention that also percolation transitions exist on Sierpinski structures [17]. Also, various hierarchical lattices different from the above fractals show phase transitions, if Ising spin are put on them; the reader is referred to [18] [19] for more literature. As the to our knowledge most recent example we mention [20] that Ising spins were also thrown into the sandpiles of Per Bak, which show self-organised criticality.

6 Other Subjects?

7 Networks

7.1 Definitions

While fractals were a big physics fashion in the 1980s, networks are now a major physics research field. Solid state physics requires nice single crystals where all atoms sit on a periodic lattice. In fluids they are ordered only over shorter distances but still their forces are restricted to their neighbours. Human beings, on the other hand, form a regular lattice only rarely, e.g. in a fully occupied lecture hall. In a large crowd they behave more like a fluid. But normally each human being may have contacts with the people in neighbouring residences, with other neighbours at the work place, but also via phone or internet with people outside the range of the human voice. Thus social interactions between people should not be restricted to lattices, but
should allow for more complex networks of connections.

One may call Flory’s percolation theory of 1941 a network, and the later random graphs of Erdős and Rényi (where everybody is connected with everybody, albeit with a low probability) belong to the same “universality class” (same critical exponents) as Flory’s percolation. In Kauffman’s random Boolean network of 1969, everybody has $K$ neighbours selected randomly from the $N$ participants. Here we concentrate on two more recent network types, the small-world \[21\] and the scale-free \[22\] networks of 1998 and 1999 respectively (with a precursor paper of economics Nobel laureate Simon \[23\] from 1955).

The small-world or Watts-Strogatz networks start from a regular lattice, often only a one-dimensional chain. Then each connection of one lattice site to one of its nearest neighbours is replaced randomly, with probability $p$, by a connection to a randomly selected other site anywhere in the lattice. Thus the limits $p = 0$ and 1 correspond to regular lattices and roughly random graphs, respectively.

In this way everybody may have exactly two types of connections, to nearest neighbours and to arbitrarily far away people. This unrealistic feature of small-world networks is avoided by the scale-free networks of Barabási and Albert \[22\], defined only through topology with (normally) no geometry involved:

We start with a small set of fully connected people. Then more people join the network, one after the other. Each new member selects connections to exactly $m$ already existing members of the network. These connections are not random but follow preferential attachment: The more people have selected a person to be connected with in the past, the higher is the probability that this person is selected by the newcomer: the rich get richer, famous people attract more followers than normal people. In the standard Barabási-Albert network, this probability is proportional to the number of people who have selected that person. In this case, the average number of people who have been selected by $k$ later added members varies as $1/k^3$. A computer program is given e.g. in \[24\].

These networks can be undirected (the more widespread version) or directed (used less often.) For the undirected or symmetric case, the connections are like friendships: If A selects B as a friend, then B also has A as a friend. For directed networks, on the other hand, if A has selected B as a boss, then B does not have A as a boss, and the connection is like a one-way street. Up to $10^8$ nodes were simulated in directed scale-free networks.
We will now check for phase transitions on both directed and undirected networks.

## 7.2 Phase transitions

The Ising model in one dimension does not have a phase transition at some positive critical temperature \( T_c \). However, its small-world generalisation, i.e. the replacement of small fraction of neighbour bonds by long-range bonds, produces a positive \( T_c \) with a spontaneous magnetisation proportional to \((T_c - T)^\beta\), and a \( \beta \) smaller than the 1/8 of two dimensions [25].

The Solomon network is a variant of the small-world network: Each person has one neighbourhood corresponding to the workplace and another neighborhood corresponding to the home [26]. It was suggested and simulated by physicists Solomon and Malarz, respectively, before Edmonds [27] criticised physicists for not having enough “models which explicitly include actions and effects within a physical space as well as communication and action within a social space.” Even in one dimension a spontaneous magnetisation was found.

On Barabási-Albert (scale-free) networks, Ising models were found [28] for small \( m \) and millions of spins to have a spontaneous magnetisation for temperatures below some critical temperature \( T_c \) which increases logarithmically with the number \( N \) of spins: \( k_B T_c / J \approx 2.6 \ln(N) \) for \( m = 5 \).

Here we had undirected networks with symmetric couplings between spins: actio = –reactio, as required by Newton. Ising spins on directed networks, on the other hand, have no well-defined total energy, though each single spin may be influenced as usual by its \( m \) neighbour spins. If in an isolated pair of spins \( i \) and \( k \) we have a directed interaction in the sense that spin \( k \) tries to align spin \( i \) into the direction of spin \( k \), while \( i \) has no influence on \( k \), then we have a perpetuum mobile: Starting with the two spins antiparallel, we first flip \( i \) into the direction of \( k \), which gives us an energy \( J \). Then we flip spin \( k \) which does not change the energy. Then we repeat again and again these two spin flips, and gain an energy \( J \) for each pair of flips: too nice to be true. The violations of Newton’s symmetry requirement makes this directed network applicable to social interactions between humans, but not to forces between particles in physics.

On this directed Barabási-Albert network, the ferromagnetic Ising spins gave no spontaneous magnetization, but the time after which the magnetisation becomes zero (starting from unity) becomes very long at low temper-
atures, following an Arrhenius law \[29\]: time proportional to \(\exp(\text{const}/T)\). Also on a directed lattice such Arrhenius behavior was seen while for directed random graphs and for directed small-world lattices a spontaneous magnetisation was found \[29\]. A theoretical understanding for these directed cases is largely lacking.

Better understood is the percolative phase transition on scale-free networks (see end of Sec.2 for definition of percolation). If a fraction \(1 - p\) of the connections in an undirected Barabási-Albert network is cut randomly, does the remaining fraction \(p\) keep most of the network together? It does, for large enough networks, since the percolation threshold \(p_c\) below which no large connected cluster survives, goes to zero as \(1/\log(N)\) where \(N\) counts the number of nodes in the network. This explains why inspite of the unreliability of computer connections, the internet still allows most computers to reach most other computers in the word: If one link is broken, some other link may help even though it may be slower \[22\].

### 8 Future Directions

We reviewed here a few phase transitions, and ignored many others. At present most interesting for future research seem to be the directed networks, since they have been investigated with methods from computational physics even though they are not part of usual physics, not having a global energy. A theoretical (i.e. not numerical) understanding would help.

### References

[1] Primary literature:

   Cont R, Bouchaud J-P (2000) Macroeconomic Dynamics 4: 170

[2] Schelling TC (1971) J. Math. Sociol. 1: 143

[3] Sznajd-Weron K, and Sznajd J (2000) Int. J. Mod. Phys. C 11: 1157

[4] Planned article by D. Stauffer for this encyclopedia (Opinion dynamics and sociophysics)

[5] Ben-Avraham D, Havlin S (1982) J. Phys. A 15: L 691
[6] Gefen Y, Aharony A, Alexander S (1983) Phys. Rev. Lett. 50: 77
[7] Kutner R, Kehr K (1983) Phil. Mag. A 48: 199
[8] Brandt WW (1975) J. Chem. Phys. 63: 5162
[9] de Gennes PG (1976) La Recherche 7: 916
[10] Frey E, Kroy K (2005) Ann. Physik 14: 20
[11] Johansen A, Sornette D (1999) Int. J. Mod. Phys. C 10: 563
[12] Kirsch A (1999) Int. J. Mod. Phys. C 10: 753
[13] Dhar D, Stauffer D (1998) Int. J. Mod. Phys. C 9: 349
[14] Berger N, Ganten N, Peres Y (2003) Probab. Theory Relat. Fields 126: 221
[15] Gefen Y, Mandelbrot BB, Aharony A (1980) Phys. Rev. Lett. 45: 855
[16] Bab MA, Fabricius G, Albano EV (2005) Phys. Rev. E 71: 036139
[17] Monceau P, Hsiao PY (2004) Phys. Lett. A 332: 310
[18] Hinczewski M, Berker AN (2006) Phys. Rev. E 73: 066126
[19] Rozenfeld HD, Ben-Abraham D (2007) cond-mat/0703155 on arXiv.org, submitted to Phys. Rev. E
[20] Koza Z, Ausloos M (2007) Physica A 375: 199
[21] Watts, D.J., Strogatz SH (1998) Nature 393: 440
[22] Albert R, Barabási AL (2002) Rev. Mod. Phys. 74: 47; Boccalettia S, Latora V, Moreno Y, Chavez M, Hwang D-U (2006) Phys. Repts. 424: 175
[23] Simon HA (1955) Biometrika 42: 425
[24] Stauffer D, Moss de Oliveira S, de Oliveira PMC, Sá Martins JS (2006) Biology, Sociology, Geology by Computational Physicists, Elsevier, Amsterdam
[25] Barrat A, Weigt M (2000) Eur. Phys. J. B 13: 547; Gitterman M. (2000), J. Phys. A 33: 8373; Pękalski A., Phys. Rev. E 64: 057104

[26] Malarz K (2003) Int. J. Mod. Phys. C 14: 561

[27] Edmonds B (2006) p. 195 in: Billari FC, Fent T, Prsakwetz A, Scheffran J (2006) Agent-based computational modelling, Physica-Verlag, Heidelberg

[28] Aleksiejuk A, Hołyst JA, Stauffer D (2006) Physica A 310: 260; Dorogovtsev SN, Goltsev AV, Mendes JFF (2002) Phys. Rev. E 66, 016104; Bianconi G (2002) Phys. Lett. A 303: 166

[29] Sánchez AD, López JM, Rodríguez MA (2002) Phys. Rev. Letters 88: 048701, Sumour MA, Shabat, MM (2005) Int. J. Mod. Phys. C 16: 585; Sumour MA, Shabat MM, Stauffer D (2006) Islamic University Journal (Gaza) 14: 209; Lima FWS, Stauffer D (2006), Physica A 359: 423; Sumour MA, El-Astal AH, Lima FWS, Shabat MM, Khalil HM (2007) Int. J. Mod. Phys. C 18: 53; Lima FWS (2007) Physica A preprint.

[30] **Books and Reviews:**

Bunde A, Havlin S (1996) Fractals and Disordered Systems, Springer, Berlin

[31] Havlin S, Ben Avraham D 1987/2002) Adv. Phys. 36, 395 and 51, 187.