Structure Below the Growing Surface

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Abstract. – In recent years there has been a growing interest in the statistical properties of surfaces growing under deposition of material. Yet it is clear that a theory describing the evolution of a surface should at the same time describe the properties of the bulk buried underneath. Clearly, the structure of the bulk is relevant for many practical purposes, such as the transport of electric current in devices, transport of fluids in geological formations and stress transmission in granular systems. The present paper demonstrates explicitly how models describing deposition can provide us with information on the structure of the bulk. Comparison of an analytic model with a simulation of a discrete growth model reveals an interesting long range tail in the density-density correlation in the direction of deposition.

The study of the statistical properties of surfaces growing under the deposition of material has attracted many researchers over the last two decades. The systems under consideration vary from heaps of sand or other assemblies of granular matter, to devices manufactured by the bombardment of atoms on a growing target. The theoretical description of such systems is given by a number of discrete and continuous models that belong mainly to three categories. The first is the Edwards Wilkinson category \cite{1} which was constructed to describe a situation of slow deposition under gravity where each deposited particle has the time to find its lowest possible gravitational potential energy in the presence of the existing surface. The second category is that of KPZ \cite{2} in which lateral growth is important. This can be a result of sticking or just the geometry of growth perpendicular to the surface as explained in ref. \cite{2}. The third category is the MBE (Molecular Beam Epitaxy) \cite{3} which was constructed to describe processes of device fabrication in which the physics should produce under a wide range of parameters flat surfaces. The focus of interest in those studies was the statistical characterization of the growing surface. This is achieved by obtaining the roughness exponent of the steady state surface, the growth exponent \cite{4–20} and the scaling functions associated with the steady state evolution of the surface \cite{21–24}. It is easy to envisage many practical applications
for which the fluctuations in the steady state surface are relevant and this was always one of
the motivations for the intensive study of surface growth. Yet it is obvious that the internal
structure of the material below the surface may be of much more practical importance as this
determines the mechanical properties of the system, generated by deposition, such as a heap
of granular matter in which it will affect stress transmission. In electrical devices obtained by
deposition, the structure of the bulk will obviously affect the transmission of electric current
which will determine the functionality of the device. Certain geological formations relevant
to the oil industry are also generated by deposition. The structure of the bulk determines the
important flow properties through the formation.

This letter has two goals. The first and more important goal is to draw attention to the
fact that a theory which describes the evolution of the upper surface of a system growing
under the deposition of material, should simultaneously be able to predict the structure of
the bulk below that surface. The physical reason for our statement is the fact that serious
rearrangement does not usually take place in the deeper layers below the surface.

The second goal is to explicitly demonstrate how the structure of bulk below the surface
can be obtained for two growth models, the continuous analytical KPZ system and the discrete
numerical ballistic deposition (BD) system. Note, that the main reason for treating the two
models mentioned above, is that both possesses as will be shown in the following share the
common realistic physical property that the growth process is accompanied by the embedding
of voids below the evolving surface (this property is not shared by other very successful discrete
numerical systems such as SOS [25] and RSOS [5] which do not allow for the generation of
voids below the surface.) The second reason for treating these two systems is that they are,
widely known and simple enough to demonstrate the connection between the system describing
the evolution of the surface and the structure of the bulk below it.

To clarify our ideas consider first the Edwards Wilkinson equation for the local height of
the surface $h(r, t)$,

$$\frac{\partial h}{\partial t} = \nu \nabla^2 h + R(r, t) ,$$

where $R(r, t)$ is the local rate of deposition of material given by

$$R(r, t) = R_0 + \eta(r, t)$$

and $\eta(r, t)$ is a noise term that has zero average and its correlations are usually taken to be

$$\langle \eta(r, t) \eta(r', t') \rangle = D \Delta(r - r') \delta(t - t'),$$

where $\Delta$ is a short range function. The constant position and time independent rate $R_0$ is tra-
ditionally deleted, because it has no effect on the shape of the surface. For our purpose though,
we have to keep it. To be precise note that the deposition rate is related to the local number
of particles landing per unit area per unit time, $n(r, t)$, by the relation

$$R(r, t) = \Omega n(r, t)$$

where $\Omega$ is the effective volume taken by each landing particle. Now particles land and rear-
range on the surface to minimize their potential (gravitational) energy. This rearrangement
is described by the first term on the right hand side of eq. (1). Dividing $h(r, t)$ by $\Omega$ we
obtain the total area density of particles at the point $r$ at time $t$. It is clear that whatever the
dynamical picture of surface evolution is, the total number of particles has to be conserved
and indeed eq. (1) trivially conserves the number of particles. Actually it does more than
that. It conserves the volume occupied by the particles. The picture is thus the following.
The landing particles form a compact structure and then rearrange on the surface preserving
the compact structure beneath it. If the particles are cubes, as in some discrete models, there
remain no voids among the particles. Other shapes of particles must result, of course, in voids
among them but the structure is expected to be either ordered or random close packing. Our
following discussion is not concerned with that compact structure. We focus rather on deviations
from that structure involving larger voids in the system. The physical reasons for such
voids include sticking and the geometry of growth normal to the surface, that are described within the KPZ category. This incorporates voids into the structure in a most natural way. To visualize it, consider the following version of ballistic deposition. In this model a particle falls vertically and sticks to the first site along its trajectory that has an occupied nearest neighbor. The particle is not allowed to stick to a diagonal neighbor. The last particle to be deposited is shaded in Fig. (1) that demonstrates how voids are created. A void, once created, does not disappear.

In the following we will discuss the problem of structure below the evolving surface from two different points of view. The first is to show that within the continuous KPZ description of the evolving surface, it is possible to obtain the density-density correlation function of the material below the surface in terms of height correlations related to the evolution of the surface. The second is to obtain the density-density correlations directly from a one dimensional BD simulation.

Consider next the KPZ equation

$$\frac{\partial h}{\partial t} = \nu \nabla^2 h + \lambda (\nabla h)^2 + R(r, t).$$

(3)

The origin of the non linear term in the above equation is the fact that growth is perpendicular to the surface. It is clear that if a particle lands on the surface an outcrop perpendicular to the surface is generated. The overhang screens the area below it and prevents more descending particles to fill the void generated by that particle just as in the discrete BD version depicted in Fig. (1). The KPZ equation must be thus viewed not as an equation for a single valued height function, because the height function is not really single valued, but rather an equation for the height envelope function below which there is an abundance of voids. Can we see the existence of those voids from the KPZ equation itself? From the relation \( R(r, t) = \Omega n(r, t) \) it is clear that the total "deposited volume" is given by \( \int dr dt R(r, t) \) (assuming we start deposition on a flat surface at time \( t = 0 \)). From eq. (3) on the other hand it is clear that the total volume below the envelop surface is larger than that by \( \lambda \int dr dt (\nabla h (r, t))^2 \) which must be the volume occupied by voids. The average of the integrand is known in the literature as the excess velocity and its meaning is that the average height growth faster than expected from the rate of descending material and this is just due to voids being incorporated into the structure [11]. Suppose we could identify that part of the local rate of increase in the height, \( \left[ \frac{\partial h(r, t)}{\partial t} \right]_v \), which is a result of void creation, then the local density \( \rho(r, z) \) at a point, through which the surface passed at time \( t \), is given by

$$\rho (r, z) = \rho_0 \left\{ \frac{\partial h (r, t)}{\partial t} - \left[ \frac{\partial h (r, t)}{\partial t} \right]_v \right\} / \frac{\partial h (r, t)}{\partial t},$$

(4)
where $\rho_0$ is the constant density that would have existed in the corresponding Edwards-Wilkinson system.

The left hand side of the above gives $\rho$ as a function of $z$ but on the right hand side we have functions of $t$. The next step must thus be to connect between the perpendicular coordinate $z$ and the passage time of the surface $t$. This relation is readily obtained by noting that the height of the surface at the point $r$ and time $t$ is given by

$$h(r, t) = R_0 t + \delta h(r, t),$$

(5)

where $\delta h(r, t)$ is the usual variable describing the width of the surface. This quantity can attain values of the order of a positive power in the lateral size of the system. Nevertheless, if we wait for long enough times, that make $R_0 t$, to be of the order of the lateral size, the second term on the right hand side of eq. (5) is negligible compared to the first. The relation can be thus iterated in the following manner

$$t = z/R_0 - \delta h(r, z/R_0 - \delta h(r, z/R_0 \ldots)),$$

(6)

Within the leading approximation to eq. (6) eq. (4) is also simplified and reads

$$\rho(r, z) = \rho_0 \left\{ 1 - \frac{[\partial h(r, t)]}{\partial t} \bigg|_{v,t=z/R_0} / R_0 \right\}.$$

(7)

We identify next

$$\frac{[\partial h(r, t)]}{\partial t} = \lambda (\nabla h(r, t))^2,$$

(8)

since this non-linear term is the term responsible for the "excess velocity" and thus for the incorporation of voids. Eqs. (7) and (8) will be used now to obtain the average density $\bar{\rho}$ and the density-density correlations in the system

$$\bar{\rho} = \rho_0 \left[ 1 - \lambda \left( \langle \nabla h \rangle^2 \right) / R_0 \right].$$

(9)

Now, assuming that the steady state structure factor is given by $\langle h_{q,h_{-q}} \rangle = Aq^{-\Gamma}$ for $q \leq q_0$ (where $q_0$ is the high momentum cut-off, corresponding to the size of landing particles) and zero above it, the average density is given by

$$\bar{\rho} = \rho_0 \left[ 1 - \lambda S_d q_0^{d+1-\Gamma} / (d + 1 - \Gamma) R_0 \right],$$

(10)

where $d+1$ is the dimension of space and $S_d$ is the surface area of a unit sphere in $d$ dimensions. The density-density correlations involve one non-trivial correlation

$$\Psi(r, t) = \langle |\nabla h(r, t)|^2 [\nabla h(0, 0)]^2 \rangle - \left[ \langle (\nabla h)^2 \rangle \right]^2.$$

(11)

To evaluate it we calculate it to the lowest order in the frequency dependent structure factor $\Phi(q, \omega) = \langle h_{q,\omega} h_{-q,-\omega} \rangle$. Standard manipulation leads to the expression

$$\Psi(r, t) = -\frac{2}{(2\pi)^2 A^2} \int dldl' (1-l')^2 l^{-\Gamma} l'^{-\Gamma} f(\omega_l t) f(\omega_{l'} t) \exp[i(l+l') \cdot r],$$

(12)

where $\omega_l = Bl^2$ is the typical frequency related to the decay of a disturbance of wave vector $l$ and the scaling function $f(u)$ is related to the time dependent structure factor $\Phi(q, t) = (2\pi)^{-1} \int d\omega \Phi(q, \omega) e^{i\omega t}$ by

$$\Phi(q, t) = Aq^{-\Gamma} f(\omega_q t).$$

(13)
Clearly, the calculation of higher orders in the iteration may be rather tedious but they are straightforward as described in refs. [13, 14]. From our point of view, however, the important statement is that density-density correlations below the growing surface are related to various height correlations which are natural objects of study in the traditional research of surface growth. It has to be carried in mind, though, that to use the correlation \( \Psi (r, t) \) for our purpose by relating the time difference to the difference in height, we must have \( R_0 t \gg |\delta h (r, 0)| , |\delta h (0, t)| \).

We will work out as an example the one-dimensional case and compare it with the simulation of the one-dimensional ballistic deposition. The density-density correlation function \( g(x, y) = \langle \rho (x) \rho (y) \rangle - \bar{\rho}^2 \) is related to \( \Psi \) in the one-dimensional case by

\[
g(x, y) = \left( \frac{\lambda}{R_0} \right)^2 \bar{\rho}^2 \Psi \left( x, \frac{y}{R_0} \right). \tag{14}\]

Using the scaling form for the 1D time dependent structure factor [11], \( \Phi (q, t) = \frac{D}{q^2} f (B t) \) (where \( D \) is the noise amplitude, \( \nu \) the diffusion coefficient and \( B \) a numerical constant which cannot be determined analytically) we obtain

\[
\Psi (x, t) = - \frac{8 D}{9 \pi^2 \nu^2 B^{4/3}} \left\{ \int_0^\infty \frac{1}{\tau^{1/3}} f(\tau) \cos \left\{ \left( \frac{\tau}{B t} \right)^{2/3} x \right\} d\tau \right\}^2. \tag{15}\]

This form is based on the scaling form and is therefore correct for large \( t \), so that for large \( y \) we obtain a power law behavior for

\[
g(0, y) = - \left( \frac{\lambda}{R_0} \right)^2 \bar{\rho}^2 \frac{8 D}{9 \pi^2 \nu^2 B^{4/3}} \left\{ \int_0^\infty \frac{1}{\tau^{1/3}} f(\tau) d\tau \right\}^2 \propto - y^{-4/3}. \tag{16}\]

It is interesting now to discuss briefly the case \( \lambda < 0 \). The parameters \( R_0 \) and \( \lambda \) corresponding to our physical picture are positive, of course. The situation with positive \( R_0 \) and negative \( \lambda \), which can be handled formally by our equations, has no easy natural interpretation. Certainly this is not related to the RSOS models, which are supposed to correspond to negative \( \lambda \) but produce compact structures.

Let us turn now to the density-density correlations obtained from a one-dimensional model of ballistic deposition. The model used is the nearest-neighbor ballistic deposition (NNBD) model, on a lattice with \( L = 1024 \). At each time step a column \( i \) is picked at random. A particle (just one particle!) falls vertically and sticks to the first site along its trajectory that has an occupied nearest neighbor as shown in Fig. 1. We found it simpler not to use periodic boundary conditions. Fig. 2 presents the structure obtained after deposition.

The alternating shades of grey correspond to particles deposited within different time intervals. The voids are the white regions. This figure corresponds to a lateral size of 512 sites.

The long-range correlations obtained in the analytical calculation above can be already observed qualitatively in the plot of the cluster generated by the simulation (Fig. 2), where some of the voids tend to extend upward for a long distance. In order to quantify this, we obtain the correlation \( g(0, y) \) from the simulation. We define the correlation function

\[
g(x, y) = \frac{1}{N} \sum \langle \rho (r) \rho (r') \rangle - \bar{\rho}^2, \tag{17}\]

where \( x \) is in the lateral direction, \( y \) is in the perpendicular (growth) direction, \( r = (X, Y) \), \( r' = (X', Y') \) such that \( |X - X'| = x \) and \( |Y - Y'| = y \). The number of such pairs \( (r, r') \) is \( N \),
Fig. 2 – A BD cluster obtained by depositing 100,000 particles on a substrate of size $L = 512$. A time step is defined by a deposition of a single particle. The different shadings correspond to different time intervals each corresponding to the deposition of 10,000 particles.

The average is over realizations of the randomness and $\bar{\rho}$ is the average density. Practically we take the average to be the average over runs. In this work we focused on correlations in the $y$ (growth) direction since they exhibit nontrivial features such as algebraic tails. We depict $g(0, y)$ in Fig. 3. The total number of pairs that goes into the evaluation of $g(0, y)$, which is the product of the number of pairs in one run times the number of runs is $2.4 \times 10^{10}$.

We depict the log of the absolute value correlation function, because the correlation function drops fast such that it is difficult to observe certain features. For example, it is difficult to see that the correlations become negative. When we depict the logarithm of the absolute value we see the point where the function changes sign as a kink. (Because the $y$-axis is a discrete coordinate the correlations do not become zero but go from positive values where they are decreasing to negative value where they are increasing). The correlation drops very fast and may seem to be of short range. The analytical prediction though suggests that we should examine more carefully the tail of $g(0, y)$. In Fig. 3(b) we depict a log-log plot of this tail, which reveals a power-law behavior of the tail, namely $g(0, y) \propto -y^{-1.301}$. We see good agreement the between the numerical result given above and the analytical one. Both yield a negative tail decreasing in size as a power law with exponents that are quite close (1.3 vs. 1.333).

Fig. 3 – (a) The log of the correlation function in the growth direction. (b) A log-log plot of the tail.
We have chosen here to concentrate on the density-density correlation function, because it is the most obvious and traditional characterization of the structure. It is clear that for specific applications like stress transmission or transmission of electrical currents we might be interested in other attributes of the structure. The main point we make in this letter is that the same analytical and numerical techniques used for the study of the growth of surfaces can be used in the study of material structure below the surface. Once this point is realized other properties of the structure can be readily be obtained. In addition, we have obtained correlations in the growth direction with an unexpected algebraic tail. If this is not just an artifact of the two models studied, it may have far reaching practical and theoretical consequences. As a possible usage we claim that the anisotropy between the growth direction and the lateral growth is a signature of the growth direction. Thus, having at hand a specimen of a sedimentary rock, for example, our result suggest that one can determine its original growth direction by determining the principal direction at which an algebraic tail in the structure factor appears.

We hope that the present work will trigger more research in this direction, both experimentally, and by introduction of more realistic discrete models of deposition.

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