Exact solution of two friendly walks above a sticky wall with single and double interactions

[†]Aleks Owczarek, [‡]Andrew Rechnitzer, and [‡]Thomas Wong

[†]MASCOS and Department of Mathematics and Statistics, The University of Melbourne

[‡] Department of Mathematics and Statistics, University of British Columbia

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DIRECTED WALKS

- Exact solutions of single and multiple directed walk models
- Recurrence and functional equation
- Rational, algebraic, Differentially-finite and non D-finite solutions
- Multiple walks: Bethe Ansatz & Lindström-Gessel-Viennot
- LGV Lemma: multiple walks = determinant of single walks
- LGV problems result in generating functions that are D-finite

SOLVING FUNCTIONAL EQUATIONS

- Functional equation for an expanded generating function
- Uses an extra catalytic variable
- Answer is a 'boundary' value
- Fix catalytic variable \rightarrow 'bulk' term disappears (Kernel method)
- Obstinate kernel method: multiple values of catalytic variable
- See Bousquet-Mélou *Math. and Comp. Sci 2* (2002)), Bousquet-Mélou, Mishna *Contemp. Math.* **520** (2010)
- Solutions are not always D-finite

POLYMER ADSORPTION

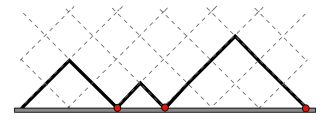
The physical motivation is the adsorption phase transition

- Second order phase transition with jump in specific heat
- Crossover exponent $\phi = 1/2$ for directed walks and SAW
- Order parameter is density of visits to surface by the polymer

ADSORPTION: ONE DIRECTED WALK

Exact solution and analysis of single and multiple directed walk models exist

- Single Dyck path in a half space
- Energy $-\varepsilon_a$ for each time (number m_a) it visits the surface
- Boltzmann weight $a = e^{\varepsilon_a/k_BT}$



ADSORPTION: ONE DIRECTED WALK

A complete solution exists and the generating function is algebraic

Consider the density of visits

$$\mathcal{A} = \lim_{n \to \infty} \frac{\langle m_a \rangle}{n}$$

There exists a phase transition at a temperature T_a given by a = 2:

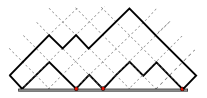
- For $T > T_a$ the walk moves away entropically and A = 0
- For $T < T_a$ the walk is adsorbed onto the surface and A > 0

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Solution and analysis

Conclusions

VESICLE ADSORPTION



- Exact solution of two directed walks joined making a simple "vesicle" (Brak, Essam, Owczarek , J. Stat. Phys. 93, 155 (1998))
- Vesicles with interactions for visits of the *bottom* walk to height 0 and height 1 (Lonsdale, Brak, Essam, Owczarek, Rechnitzer, *J. Phys. A.: Math. and Theor.* **42** 1, (2009).)

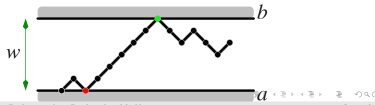
Single second order transition — similar to the single walk adsorption transition

MORE MOTIVATION: POLYMERS IN A SLIT

Paths in a slit

- A directed path of length *n* in a slit of width *w*.
- Boltzmann weights *a* and *b* for interactions with lines (walls).
- Phase diagram depends on order of limits
- Brak, Owczarek, Rechnitzer and Whittington J. Phys. A, **38** (2005) 4309–4325
- Subsequent works on undirected SAW
- Different regions of parameter space yield different effective types of force the polymer has on the walls of the slit

• simple hyperbolic "zero force curve"



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RING POLYMERS IN A SLIT

- Our motivation is a recent Monte Carlo study of ring polymers in a slit
- Here both sides of the polygon interact with the surfaces of the slit Alvarez, Janse van Rensburg, Soteros and Whittington *J. Phys. A.: Math. and Theor.* **41**, 185004 (2008)
- Novel zero force curve

(Our Model)

Directed vesicle where both walks can interact with a single surface

Model

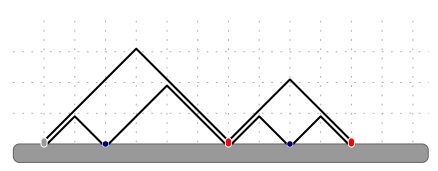


Figure: Two directed walks with single and "double" visits to the the surface.

- energy $-\varepsilon_a$ for visits of the bottom walk only (single visits) to the wall,
- energy $-\varepsilon_d$ when both walks visit a site on the wall (double visits)

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- number of *single visits* to the wall will be denoted m_a ,
- number of *double visits* will be denoted *m*_d.

The partition function:

$$Z_n(a,d) = \sum_{\widehat{\varphi} \ni |\widehat{\varphi}| = n} e^{(m_a(\widehat{\varphi}) \cdot \varepsilon_a + m_d(\widehat{\varphi}) \cdot \varepsilon_d)/k_B T}$$

where $a = e^{\varepsilon_a/k_BT}$ and $d = e^{\varepsilon_d/k_BT}$.

The thermodynamic reduced free energy:

$$\kappa(a,d) = \lim_{n\to\infty} n^{-1} \log \left(Z_n(a,d) \right).$$

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GENERATING FUNCTION

To find the free energy we will instead solve for the generating function

$$G(a,d;z) = \sum_{n=0}^{\infty} Z_n(a,d) z^n.$$

The radius of convergence of the generating function $z_c(a, d)$ is directly related to the free energy via

$$\kappa(a,d) = \log(z_c(a,d)^{-1}).$$

Two order parameters:

$$\mathcal{A}(a,d) = \lim_{n \to \infty} \frac{\langle m_a \rangle}{n}$$
 and $\mathcal{D}(a,d) = \lim_{n \to \infty} \frac{\langle m_d \rangle}{n}$,

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FUNCTIONAL EQUATION

We consider walks φ *in the larger set, where each walk can end at any possible height.*

The expanded generating function

$$F(r,s;z) \equiv F(r,s) = \sum_{\varphi \in \Omega} z^{|\varphi|} r^{\lfloor \varphi \rfloor} s^{\lceil \varphi \rceil/2} a^{m_a(\varphi)} d^{m_d(\varphi)},$$

where

- *z* is conjugate to the length $|\varphi|$ of the walk,
- *r* is conjugate to the distance $\lfloor \varphi \rfloor$ of the bottom walk from the wall and
- *s* is conjugate to *half* the distance [φ] between the final vertices of the two walks.

and we recover G(a, d; z) = F(0, 0).

FUNCTIONAL EQUATION

Consider adding steps onto the ends of the two walks

This gives the following functional equation

$$F(r,s) = 1 + z \left(r + \frac{1}{r} + \frac{s}{r} + \frac{r}{s}\right) \cdot F(r,s) - z \left(\frac{1}{r} + \frac{s}{r}\right) \cdot [r^{0}]F(r,s) - z\frac{r}{s} \cdot [s^{0}]F(r,s) + z(a-1)(1+s) \cdot [r^{1}]F(r,s) + z(d-a) \cdot [r^{1}s^{0}]F(r,s).$$

Figure: Adding steps to the walks when the walks are away from the wall.

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THE KERNEL

Rewrite equation as "Bulk = Boundary"

$$K(r,s) \cdot F(r,s) = \frac{1}{d} + \left(1 - \frac{1}{a} - \frac{zs}{r} - \frac{z}{r}\right) \cdot F(0,s) - \frac{zr}{s} \cdot F(r,0) + \left(\frac{1}{a} - \frac{1}{d}\right) \cdot F(0,0)$$

where the kernel K is

$$\mathsf{K}(r,s) = \left[1 - z\left(r + \frac{1}{r} + \frac{s}{r} + \frac{r}{s}\right)\right].$$

Recall, we want F(0,0) *so we try to find values that kill the kernel*

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SYMMETRIES OF THE KERNEL

The kernel is symmetric under the following two transformations:

$$(r,s)\mapsto \left(r,\frac{r^2}{s}\right),$$
 $(r,s)\mapsto \left(\frac{s}{r},s\right)$

Transformations generate a family of 8 symmetries ('group of the walk')

$$(r,s), \left(r, \frac{r^2}{s}\right), \left(\frac{s}{r}, \frac{s}{r^2}\right), \left(\frac{r}{s}, \frac{1}{s}\right), \left(\frac{1}{r}, \frac{1}{s}\right), \left(\frac{1}{r}, \frac{s}{r^2}\right), \left(\frac{r}{s}, \frac{r^2}{s}\right), \text{ and } \left(\frac{s}{r}, s\right)$$

We make use of 4 of these which only involve positive powers of r.

This gives us four equations.

MAGIC COMBINATION

Following Bousquet-Mélou when a = 1 we form the simple alternating sum

$$Eqn1 - Eqn 2 + Eqn 3 - Eqn 4$$

- When $a \neq 1$ one needs to generalise that approach
- Multiply by rational functions,

The form of the Left-hand side of the final equation being

$$a^{2}rK(r,s)\left(sF(r,s) - \frac{r^{2}}{s}F\left(r,\frac{r^{2}}{s}\right) + \frac{Lr^{2}}{s^{2}}F\left(\frac{r}{s},\frac{r^{2}}{s}\right) - \frac{L}{s^{2}}F\left(\frac{r}{s},\frac{1}{s}\right)\right)$$

where

$$L = \frac{zas - ars + rs + zar^2}{zas - ar + r + zar^2}.$$

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EXTRACTING THE SOLUTION a = 1

 $K(r,s) \cdot (\text{linear combination of } F) =$

$$\frac{r(s-1)(s^2+s+1-r^2)}{s^2} \left(1+(d-1)F(0,0)\right) \\ -zd(1+s)sF(0,s) + \frac{zd(1+s)}{s^2}F\left(0,\frac{1}{s}\right).$$

- The kernel has two roots
- choose the one which gives a positive term power series expansion in z
- with Laurent polynomial coefficients in s:

$$\hat{r}(s;z) \equiv \hat{r} = \frac{s\left(1 - \sqrt{1 - 4\frac{(1+s)^2 z^2}{s}}\right)}{2(1+s)z} = \sum_{n>0} C_n \frac{(1+s)^{2n+1} z^{2n+1}}{s^n},$$

where $C_n = \frac{1}{n+1} {\binom{2n}{n}}$ is a Catalan number.

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EXTRACTING THE SOLUTION a = 1

- *Make the substitution* $r \mapsto \hat{r}$
- rewrite to remove z: $z = (\hat{r} + 1/\hat{r} + \hat{r}/s + s/\hat{r})^{-1}$.

Setting $r \mapsto \hat{r}$ gives

$$0 = ds^4 F(0,s) - dsF\left(0,\frac{1}{s}\right) - (s-1)(s^2 + s + 1 - \hat{r}^2)(s + \hat{r}^2)\left(1 + (d-1)F(0,0)\right)$$

Note coefficients of F(0, s) and F(0, 1/s) are independent of \hat{r} .

If we divide by equation by s — then F(0,0) is the constant term in s.

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Solution and analysis

Conclusions

SOLUTION FOR a = 1

Now extract the coefficient of s^1 :

$$0 = -\left(1 + \sum_{n=0}^{\infty} \frac{12(2n+1)}{(n+2)^2(n+3)} C_n^2 z^{2n+2}\right) \cdot (1 + (d-1)F(0,0)) - d \cdot F(0,0).$$

Solving the above when d = 1 gives

$$G(1,1;z) = 1 + \sum_{n=0}^{\infty} \frac{12(2n+1)}{(n+2)^2(n+3)} C_n^2 z^{2n+2},$$

and hence for general *d* we have

$$F(0,0) = G(1,d;z) = \frac{G(1,1;z)}{d + (1-d)G(1,1;z)}$$

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Model and Obstinate Kernel

Solution and analysis

Conclusions

a = d

Need to extract coefficients term by term in a to give

$$\begin{aligned} [a^{k}z^{2n}]F(0,0) &= \sum_{k'=0}^{k} \frac{k'(k'+1)(2+4n-k'n-2k')}{(k'-1-n)(n+1)^{2}(-2n+k')(n+2)} \binom{2n-k'}{n} \binom{2n}{n} \\ &= \frac{k(k+1)(k+2)}{(2n-k)(n+1)^{2}(n+2)} \binom{2n-k}{n} \binom{2n}{n} \end{aligned}$$

which gives us

$$G(a,a) = \sum_{n \ge 0} z^{2n} \sum_{k=0}^{n} a^k \frac{k(k+1)(k+2)}{(n+1)^2(n+2)(2n-k)} {2n \choose n} {2n-k \choose n}.$$

Agrees with Brak et al. (1998) that used LGV

One can now consider $d \neq a$:

$$G(a,d;z) = \frac{aG(a,a;z)}{d + (a-d)G(a,a;z)}.$$

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COMBINATORIAL STRUCTURE

- Combinatorial structure the underlying the functional equation.
- Breaking up our configurations into pieces between double visits gives

$$G(a,d;z) = \frac{1}{1 - dP(a;z)}$$

where P(a; z) is the generating function of so-called primitive factors.

• Rearranging this expression gives

$$P(a;z) = \frac{G(a,d;z) - 1}{dG(a,d;z)} = \frac{G(a,a;z) - 1}{aG(a,a;z)}.$$

• This allows us to calculate P(a; z) from a known expression for G(a, a; z)

PHASES

The phases determined by dominant singularity of the generating function

The singularities of G(a, d; z) *are*

- those of P(a; z) which are related to those of G(a, a; z) and
- the simple pole at 1 dP(a; z) = 0.

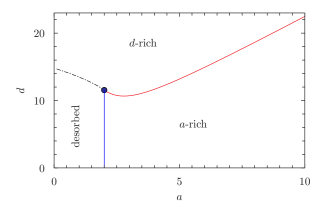
There are two singularities of G(a, a; z) giving rise to two phases:

- A desorbed phase: A = D = 0
- The bottom walk is adsorbed (an *a*-rich phase): A > 0 with D = 0

The simple pole in 1 - dP(a; z) = 0 gives rise to the third phase

• Both walks are adsorbed and this is a *d*-rich phase: D > 0, and A > 0

PHASE DIAGRAM



The first-order transition is marked with a dashed line, while the two second-order transitions are marked with solid lines. The three boundaries meet at the point $(a, d) = (a^*, d^*) = (2, 11.55...)$.

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PHASE TRANSITIONS

- The Desorbed to *a*-rich transition is
 - the standard second order adsorption transition
 - on the line a = 2 for $d < d^*$
- On the other hand the Desorbed to *d*-rich transition is first order
- While the *a*-rich to *d*-rich transition is also second order.

The other two phase boundaries are solutions to equations involving G(a, a)

The point where the three phase boundaries meet can be computed as

$$(a^*, d^*) = \left(2, \frac{16(8-3\pi)}{64-21\pi}\right)$$

Note that d^* is not algebraic.

NATURE OF THE SOLUTION

Desorbed to *d*-rich transition occurs at a value of $d_c(a)$ for a < 2. We found

$$d_c(1) = \frac{8(512 - 165\pi)}{4096 - 1305\pi}$$

which is not algebraic.

- If generating function were D-finite then $d_c(1)$ must be algebraic
- Hence generating function is not D-finite
- though it is calculated in terms of one.

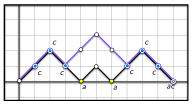
FIXED ENERGY RATIO MODEL FAMILY



- r = 2 model has *two* phase transitions as temperature changed .
- At very low temperatures the model is in a *d*-rich phase
- while at high temperatures the model is in the desorbed state.
- At intermediate temperatures the system is in an *a*-rich phase.
- Both transitions are second-order with jumps in specific heat.

CONCLUSIONS FOR DOUBLE INTERACTION MODEL

- Vesicle above a surface both sides of the vesicle can interact
- Exact solution of generating function
- Obstinate kernel method with a minor generalisation
- Solution is not D-finite LGV lemma does not apply directly
- There are two low temperature phases
- Line of first order transition and usual second order adsorption.
- Owczarek, Rechnitzer, and Wong, J. Phys. A: Math. Theor., 45 425002, (2012)
- Current work with *Rami Tabbara* on DNA unzipping model completed and submitted to *J. Phys. A: Math. Theor.*



Also work on multiple walks in slits with interactions has been completed

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