Theta-point polymers in the plane and Schramm-Loewner evolution

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We study the connection between polymers at the θ temperature on the lattice and Schramm-Loewner chains with constant step length in the continuum. The latter realize a useful algorithm for the exact sampling of tricritical polymers, where finite-chain effects are excluded. The driving function computed from the lattice model via a radial implementation of the zipper method is shown to converge to Brownian motion of diffusivity κ = 6 for large times. The distribution function of an internal portion of walk is well approximated by that obtained from Schramm-Loewner chains. The exponent of the correlation length ν and the leading correction-to-scaling exponent Δ₁ measured in the continuum are compatible with ν = 4/7 (predicted for the θ point) and Δ₁ = 72/91 (predicted for percolation). Finally, we compute the shape factor and the asphericity of the chains, finding surprising accord with the θ-point end-to-end values.

I. INTRODUCTION

Self-avoiding walks and percolation are the quintessential geometric critical phenomena. In such systems, either the statistical ensemble or the properties investigated are of a geometric nature. For instance, a self-avoiding walk (SAW) is a nearest-neighbor walk on a lattice such that each site is visited at most once; the constraint is entirely configurational. One of the most prominent characteristics of these systems is their fractal geometry, the ubiquity of which is nowadays a well-established observation [1]. Mappings exist between ordinary and geometric critical phenomena, such as the celebrated result of de Gennes [2] (proved non-perturbatively in [3]) concerning the N → 0 limit of the O(N) model. Geometric phenomena are thus interesting both per se and for their descriptive power, and also as testing grounds for ideas and tools in statistical mechanics.

In this perspective, it is perhaps not surprising that certain properties of percolation can be mapped [4] to those of an interacting version of SAW (called ISAW), in which adjacent occupied sites give a negative contribution to the energy, i.e. they attract each other, thus counteracting the opposite tendency due to the self-avoidance. ISAW are a good model for polymers in solution, as they provide an additional tunable parameter that can modulate the strength of the effective solvent-mediated interaction between monomers, on top of the repulsive excluded-volume effect. As the temperature is varied, the long-chain behavior undergoes a phase transition between an elongated SAW-like phase and a dense globular one. The transition, called the θ point, is tricritical. In fact, polymeric behavior in two spatial dimensions is diverse, and lattice models are being invented or rediscovered that display as surprising as rich a phenomenology [5] [6].

It is decisive, in the study of polymers and geometric critical phenomena, to have available the most assorted set of tools, both theoretical and numerical. Efforts and results in both areas have often proceeded on the same pathway, with enriching exchanges in both directions [7]. A new tool often comes in the form of a new model, perhaps easier to analyze or to simulate, perhaps providing deeper insight [8]. A strategy recently proposed stems from the definition of “endless” walks [9], i.e. lattice walks that can be infinitely concatenated, thus excluding any finite-chain effects. A similar property holds for another ensemble [10], based on a suitable discretization of Schramm-Loewner evolution (SLE), which gives rise to an exact-sampling algorithm for self-avoiding paths in the complex plane, in the same universality class of SAW.

Here, with these motivations in mind, we first explore the connection between SLE and θ-point polymers, showing how the two models are connected. Then we employ the aforementioned discretized SLE (such that the Euclidean length of the steps is approximately constant along the curve) to explore the connection further and to measure critical exponents and properties related to the shape of the curves.

SLE (for a nice introduction “for physicists” see [11]) is a stochastic process taking values on conformal maps in a domain. Let us fix the domain to be the unit circle D. For each time t, a random map gₜ(z) is identified as the solution to the Loewner equation

$$\partial_t g_t(z) = g_t(z) \frac{\exp(i\xi_t) + g_t(z)}{\exp(i\xi_t) - g_t(z)}, \quad g_0(z) = z,$$

where the driving function ξₜ is the one-dimensional stochastic process ξₜ = √κBₜ; κ is a parameter (playing a crucial role in the theory) and Bₜ is standard Brownian motion. The complement of the domain of gₜ, or equivalently the set of points for which gₜ does not admit a solution up to time t, is a random fractal curve, the properties of which depend on κ; notably, their fractal dimension is 1 + κ/8. The curves germinate from the point z = exp(iξ₀) and grow towards z = 0 for large times. This geometry is called radial, as opposed to the chordal one where the curve grows from a boundary point to a boundary point. By complex inversion, one can then
define curves living in the complement of $\mathbb{D}$ in the complex plane, growing towards $z = \infty$ (see Sec. [II]). SLE has been shown (either rigorously or by numerical and approximate means) to give the scaling behavior of a large number of critical models for different values of $\kappa$ [12][16]. Moreover it is the starting point for many generalisations and extensions [17][20].

SLE is a natural candidate for the continuum counterpart of polymers, as it describes non-self-crossing critical curves. Its connection with conformal field theory [21] sets a relation between the central charge $c$ and the parameter $\kappa$, namely $c = (8 - 3\kappa)(\kappa - 6)/(2\kappa)$. The only two values corresponding to $c = 0$, which is expected for critical and tricritical polymers, are $\kappa = 8/3$, which is known to describe SAW [22][23], and $\kappa = 6$, which corresponds to percolation [24] and is the obvious suspect for the $\theta$ transition. The fractal dimension, which has been calculated exactly for the latter [25], agrees with the SLE value 7/4.

Curves in SLE, as we already noted, naturally extend to infinity, and considering the evolution up to a time $t$ equates to taking a finite portion of the whole curve. Lattice polymers, on the contrary, are affected by finite-chain effects, due to their not extending to infinity; these effects are caused by configurations where the endpoint is “trapped” in a loop, which can appear at every scale at the critical point, so that finite-chain behavior persists even when the degree of polymerization $n$ (i.e. the length of the polymers) is sent to infinity in the critical limit. When endpoint distribution functions are considered, for example, the scaling variable is $\rho = r|/\xi$, where $r$ is the position of the endpoint and $\xi$ is a correlation length; both $\xi$ and $r$ scale as some power of $n$ (see Sec. [II]).

Moreover, the match between quantities on the SLE side with the corresponding properties of lattice polymers requires consideration of whether they are parameterization-independent or not, as counting the number of steps on the lattice does not in general correspond to measuring time in the continuum model. As far as SAW are concerned, these issues have been addressed [10][26]: the radial distribution of a properly chosen point along an SLE curve at $\kappa = 8/3$ matches that of an *internal* point in a lattice SAW; in other words, the polymers are “endless”, in the spirit of [9]. There are two ways of choosing the right point on the SLE side: first, one can measure the fractal variation along the curve and stop when it reaches a given value [27]; second, one can produce discrete curves (chains) so that consecutive points lie an approximately constant distance apart from each other [11]. The latter is the strategy we employ here.

The model of interacting self-avoiding walks that we are going to adopt is the standard ISAW on the square lattice. The ensemble is that of all self-avoiding nearest-neighbor $n$-step walks $\omega$, with the Gibbs measure defined by the following energy function:

$$H(\omega) = - \sum_{i=0}^{n-3} \sum_{j=i+3}^{n} \delta_{|\omega_i - \omega_j|, 1},$$  \hspace{1cm} (2)

which simply counts the number of nearest-neighbor contacts between non-bonded monomers. We will study the model at the inverse temperature corresponding to the transition between good and poor solvent, $\beta_0$. The best estimate on the square lattice is $\beta_0 = 0.6673(5)$ [28]; we will use this central value in the following.

**II. DRIVING FUNCTION**

An approximation to the driving function can be obtained by the *zipper* prescription [29]. Schematically, it amounts to applying iteratively a simple conformal map to all points of a lattice walk, mapping them in sequence to the boundary. It is usually done in the half plane, by means of vertical-slit or tilted-slit mappings, but we will apply the same strategy on the plane.

Consider the conformal map

$$\phi_t(z) = \frac{1}{2e^{\beta t}z} \left[(z + 1)^2 - 2e^\beta z \right] - (z + 1)^{1/2}\left[(z + 1)^2 - 4e^\beta z \right],$$  \hspace{1cm} (3)

which is the solution to [1] for a constant $\xi_t = 0$. The domain which gets mapped by $\phi_t$ onto the unit disc is the unit disc with a radial slit removed; the slit sticks out of $z = 1$ and is directed towards $z = 0$. The length of the slit increases with $t$ non-linearly. Let us consider the map

$$f_t(z) = \frac{1}{g_t(1/z)};$$  \hspace{1cm} (4)

it sends the exterior of the slitted disc conformally onto the exterior of the disc. The expansion around infinity of its inverse,

$$f_t^{-1}(w) = e^\beta w + 2(e^\beta - 1) + \mathcal{O}(1/w),$$  \hspace{1cm} (5)

identifies the (logarithmic) capacity of the slitted disc $\text{cap}(t) = e^\beta$, and its conformal center $c(t) = 2(e^\beta - 1)$.

Let $\{\omega_i\}$, $i = 0, \ldots, n$, be a lattice walk. Here, the lattice is embedded in the complex plane (say, with lattice spacing $a = 1$). The zipper algorithm is a prescription to find a driving function giving rise (via the Loewner equation) to an approximation of $\omega$. Operatively, at the $k$-th step one applies, to all points $\{\omega_i\}$ with $i \geq k$, the radial-slit mapping $\varphi$ that maps $\omega_k$ to the unit circle, that is

$$\varphi(\omega_k; z) = f_k(e^{-i\theta_k}z),$$  \hspace{1cm} (6)

where

$$\theta_k = \arg(\omega_k)$$  \hspace{1cm} (7)

and the time $t_k$ is obtained by solving $f_k(|\omega_k|) = 1$, i.e.,

$$t_k = \log \left(\frac{|\omega_k| + 1}{4|\omega_k|}\right).$$  \hspace{1cm} (8)
Now, $\theta_k$ is a compact variable, but since the driving function for SLE is supposed to be continuous then the interval $[0, 2\pi]$ can be unwrapped univocally onto the whole real line to obtain the approximate driving function $\xi_k$. Operatively, we set $\xi_0 = \theta_0 = w_0 = 0$ and compute $\delta_k = \theta_k - \theta_{k-1}$ at each step; whenever $|\delta_k| > \pi$ we increase or decrease the winding number $w_k$. The driving function is then $\xi_k = \theta_k + 2\pi w_k$.

We generated $\theta$-point walks in the full plane with increasing lengths ($n = 5000, 10000, 20000, 40000$) and obtained $10^4$ samples for each $n$. The method used is the extended reptation algorithm [30, 31]; remark that very efficient algorithms are available for the non-interacting SAW [32, 33], but they are expected to perform much less efficiently for the ISAW. In order to reduce the severity of finite-chain effects, we restricted the analysis of the driving function to the first $n/10$ steps (we obtain similar results with $n/20$ steps; an independent check that corrections in this regime are small is given in Fig. 3 for the radial distribution functions, see Sec. III). The function $\xi_t$ computed by the zipper algorithm turns out to be compatible with a Brownian motion with the expected diffusivity $\kappa = 6$, after an initial transient where deviations from this behavior dominate. The plots in Fig. 1 show how $\langle \xi_t^2 \rangle$ depends on $t$ (the average is over all samples generated). The expected behavior $\langle \xi_t^2 \rangle \sim \kappa t$ is recovered for large times, increasingly well in $n$. The initial transient displays an effectively higher diffusivity (around the critical value $\kappa = 8$, where curves become space-filling) followed by a lower one, which slowly relaxes towards $\kappa = 6$. We note that in the whole plane (contrary to the half-plane case) equal-length steps in the walk contribute a smaller and smaller capacity as they get further away from the origin. Therefore, although the initial transient is present up to times $t \approx 2$, it really regards only the first few walk steps (approximately 100). The falloffs at large $t$ are due to the fact that not all generated walks reach such large capacity; those that do are then biased to be more straight, and thus their driving function fluctuates less. Table I shows numerical values of $\kappa$ obtained by linear fits with several cutoffs; the upper cutoffs are chosen so as to exclude the aforementioned biased region.

Table I: Fit results for the curves in Fig. 1 with several lower cutoffs on time; the upper cutoffs are fixed by simple considerations related to biasing (see the text). The closest value to the expected $\kappa = 6$ is obtained for the longest walks and the highest cutoff.
Table II: The goodness-of-fit $p$-value obtained by the standard \( \chi^2 \) test. All $p$-values for the higher cutoff $t_{\text{min}} = 2.5$ are above the stringent significance level 0.05.

\[
\begin{array}{|c|c|c|c|c|}
\hline
n & t_{\text{min}} & t_{\text{max}} & p\text{-value} \\
\hline
5000 & 2 & 2.7 & 0.67 \\
10000 & 2 & 3.1 & 0.001 \\
20000 & 2 & 3.5 & 0.40 \\
40000 & 2 & 3.9 & 0.004 \\
5000 & 2.5 & 2.7 & 0.90 \\
10000 & 2.5 & 3.1 & 0.12 \\
20000 & 2.5 & 3.5 & 0.60 \\
40000 & 2.5 & 3.9 & 0.06 \\
\hline
\end{array}
\]

\( \kappa = 6 \), we expect

\[
\frac{\xi_t}{\sqrt{\kappa t}} \sim \mathcal{N}(0, 1), \quad (9)
\]

where \( \mathcal{N}(0, 1) \) is the normal distribution of mean 0 and variance 1. Since the foregoing analysis shows that deviations from the expected diffusivity are present for $t \leq 2$ we compute \( \xi_t/\sqrt{\kappa t} \) by using only data with $t > 2$. Figure 2 shows the results, superimposed on the Gaussian $p(x) = \exp\left(-x^2/2\right)/(2\pi)^{1/2}$. As a quantitative test, we compute the Pearson’s \( \chi^2 \) $p$-values for several cutoffs in $t$ (Tab. II); after the transient, the driving function is solidly Gaussian.

**III. DISTRIBUTION FUNCTIONS**

The opposite procedure to that of the previous section can be performed, in such a way as to generate curves \( \gamma \) in the complex plane. Instead of absorbing the points of the walk into the disc by use of the incremental map \( f_t \), one uses \( f_t^{-1} \) to grow them. The driving function here is fixed to (a suitable discretization of) Brownian motion, and the collection of points \( \{\gamma_i\} \) is what the algorithm produces. As a prescription, we approximate \( B_t \) by a piecewise constant function, because then \( \phi \) is the solution to (1) in each one of the intervals where the driving function is constant, as already noted. Hence, composition of the maps \( f_t f_t^{-1} \), where \( \delta t \) is the time step, intertwined with rotations by \( \theta \), where \( \theta \) follows a random walk, iteratively grows the set \( \{\gamma_i\} \). This is the standard way that (radial) SLE is simulated, through the so-called backward evolution; for details, see [10, 11].

We need a way of directly comparing distribution functions of lattice \( \theta \)-point walks and paths obtained by such a discretization of (1). In order to do so, the parameterizations in the two models must coincide. Schramm-Loewner paths come with a natural proper time, which is the one imposed by an exponentially increasing logarithmic capacity (a linearly increasing half-plane capacity in the chordal geometry), and lattice walks have their own, induced by the requirement that consecutive points belong to nearest neighbors. So what we need is a way of generating continuum walks \( \gamma \) (on the SLE side) such that the Euclidean lengths of their steps \( |\gamma_{i+1} - \gamma_i| \) be constant along the chain, at least approximately. Such a task encounters some technical difficulties, which nonetheless can be overcome. The method we employ, first proposed in [10], rescales each step in the iterated-map approach by an appropriate factor, obtained by tracking the evolution of the Jacobian of the map in an affordable way. We do not describe the details here, and the reader is referred to the original article.

Another obstruction to the program of matching the distribution functions of SLE and tricritical polymers is related to the fact that the full plane and the exterior of the disc are not conformally equivalent. When dealing with self-avoiding walks (corresponding to \( \kappa = 8/3 \)) one can take the position that conformal restriction, a special property concerning restrictions of the domain, which is valid only for that value of \( \kappa \), makes the distinction irrelevant. In fact, no problem arises for the SAW [20]. But in the case of \( \theta \)-point polymers, we are forced to take care of the domain. Given a lattice walk \( \omega \) of length \( n \), the sub-walk \( \hat{\omega} = \{\omega_i\}, i = 0, \ldots, l \), must be mapped to the unit disc by a conformal map \( \phi_{\hat{\omega}} \) (more precisely, the complement in \( \mathbb{C} \) of the piece-wise linear curve having vertices in \( \{\omega_i\} \) is mapped onto the exterior of the unit disc). Then the distribution functions of \( \{\phi_{\omega}(\omega_i)\}, i = l + 1, \ldots, m, \) are expected to match those obtained from SLE, provided the scaling limit \( m, n \to \infty \) is performed, with \( m \ll n \) (so as to probe the walk far from the endpoint) and \( (m-l) \to \infty \) (scaling limit of the subwalk). Note that this entails the evaluation of a random (i.e., realization
dependent) conformal map for each walk.

We do not embark here on such an exploration, and rather pursue an approximation. Namely, we consider the distribution functions for the internal sub-walk \( \{\omega_i\}, \) \( i = 1 + 1, \ldots, m \) itself, without applying the map \( \phi_\omega \). We will suppose that, given an \( n \)-step \( \theta \)-point walk \( \omega \), the probability that the \( l \)-th and \( m \)-th points (with \( l < m \)) are such that \( \omega_m - \omega_l = r \) has the following scaling form, when \( n, m, l \rightarrow \infty, r \rightarrow \infty \), keeping \( l/n = \lambda_1 \) and \( m/n = \lambda_2 \) fixed:

\[
P_{n,l,m}(r) \sim \frac{1}{\xi_l^2 \xi_m^2} F_{\text{ISAW}}(\rho, \lambda_1, \lambda_2),
\]

where \( \rho = |r|/\xi_l \) and \( \xi_{l,m}^2 = \langle |\omega_m - \omega_l|^2 \rangle \). (In measuring \( F_{\text{ISAW}} \) from data, lattice effects have been alleviated by a suitable averaging procedure that takes into account the number of lattice points in each annulus \( [\rho, \rho + \Delta \rho] \).

After taking this continuum limit, we want to let \( \lambda_2 \rightarrow 0 \) (and hence also \( \lambda_1 \rightarrow 0 \)), in order to exclude the finite-chain effects due to the presence of the endpoint. Such a complicated non-commutative limit is tricky to analyze numerically, so we give up a detailed quantitative study, and just show that the distribution functions for fixed \( n \) and several values of \( l \) and \( m \), calculated as in [10], collapse on the same curve, thus confirming the scaling form (see Fig. 3), deviations from the scaling behavior are apparent only for the largest values of \( m \) and \( l \), i.e., when \( \lambda_1 \) and \( \lambda_2 \) are significantly different from 0). On the SLE side, the distribution function is defined on the same lines. The probability density that the \( n \)-th point along the discrete chain \( \gamma \) is in \( z \), for \( n \rightarrow \infty \) and \( z \rightarrow \infty \), behaves as

\[
P_n(z) \sim \frac{1}{\xi_n^2} F_{\text{SLE}}(\rho),
\]

where \( \rho = z^*/\xi_n \) and \( \xi_{n}^2 = \langle \gamma_n \gamma_n^* \rangle \). This scaling form has been already verified numerically for \( \kappa = 8/3 \) [20], and here we check it for \( \kappa = 6 \); the inset in Fig. 3 shows the collapse of \( F_{\text{SLE}}(\rho) \) on the universal curve for chains of different lengths.

Finally, we compare the distribution functions \( F_{\text{SLE}}(\rho) \) and \( F_{\text{ISAW}}(\rho, \lambda_1, \lambda_2) \) by collapsing all data with \( n = 5000, 10000, 20000 \) for SLE and all data with \( m = 1000, 2000, 4000 \) for ISAW. As argued at the beginning of this section, we do not expect the two functions to be equal. Nonetheless, effects due to the domain are supposed to be irrelevant in the large-\( \rho \) regime, so \( F_{\text{ISAW}} \) and \( F_{\text{SLE}} \) should agree in this limit. Figure 4 shows qualitatively that this is the case, and additionally that the distribution functions only slightly disagree for small \( \rho \). Fits against the ansatz \( \log F(\rho) = c - D\rho^\delta \), inspired by the known large-distance form for \( \theta \)-point ISAW [28], yield \( \delta = 3.2(3) \) in the case of \( F_{\text{ISAW}} \) (by averaging over windows with different lower cutoffs on \( \rho \)), and \( \delta = 3.1(1) \) in the case of \( F_{\text{SLE}} \); values of \( c \) and \( D \) are compatible as well. In contrast, the endpoint distribution function and that for a single internal point, corresponding to setting \( l = 0 \) in [10], have very different shapes (Fig. 4).

\[
\langle R_{m}^2 \rangle_{m^{2\nu}} = a + \frac{a_1}{m} + \frac{a_2}{m^2} + \ldots + \frac{b_0}{m^{\Delta_1}} + \frac{b_1}{m^{\Delta_1+1}} + \ldots \tag{12}
\]

where the first terms are analytical corrections and the others have non-integer exponents. The asymptotic form [12] defines the Flory exponent \( \nu \) and the leading correction-to-scaling exponent \( \Delta_1 \). The numerical estimation of \( \Delta_1 \) is often complicated by the superposition of different terms (especially if they have discarding signs [35]) and by lattice artifacts. In the case of self-avoiding walks the SLE approach has proven interesting [10], as it displays the corrections with \( \Delta_1 = 11/16 \) that are predicted by conformal field theory [39] but not detectable for lattice self-avoiding walks [35] (not even in their endless formulation [9]). Let us fix an increment \( \delta m \) and consider \( \langle R_{m}^2 \rangle \) when \( m \) varies in steps of length \( \delta m \). We can define an effective Flory exponent \( \nu_{\text{eff}} \) as

\[
\nu_{\text{eff}}(m) = \frac{1}{2} \log \left( \frac{\langle R_{m+\delta m}^2 \rangle}{\langle R_{m}^2 \rangle} \right) \left( \frac{\log m + \delta m}{m} \right)^{-1}, \tag{13}
\]
which is simply equal to the critical exponent $\nu$ if the end-to-end radius is exactly proportional to $m^{2\nu}$. Its dependence on $m$ will highlight the corrections to scaling; in particular, if only the term with exponent $\Delta_1$ in (12) is kept, one has $\nu_{\text{eff}}(m) - \nu \propto m^{-\Delta_1}$ asymptotically.

We generated $\sim 150\,000$ independent samples of discrete SLE chains of length $n = 10\,000$ (at $\kappa = 6$), and measured $\langle R_m^2 \rangle$ for $m = j \cdot \delta m$, with $j = 1, \ldots, 40$ and $\delta m = 250$. Notice that the values obtained for different values of $m$ are not independent, so our estimates are probably affected by slight systematic errors that are difficult to quantify (the uncertainties are probably underestimated as well). Setting the ansatz $\langle R_m^2 \rangle \propto m^{2\nu}(c + m^{-\Delta_1})$ in (13) gives a form against which data can be compared. Varying both exponents $\nu$ and $\Delta_1$ in the fit is unfeasible, but Fig. 4 shows that the effective exponent converges to the expected value $\nu = 4/7 \approx 0.5714$. This is confirmed by fits of the form $\langle R_m^2 \rangle \propto m^{2\nu}$ for the radius, performed in windows $m \in [m_{\text{min}}, 10\,000]$ for increasing values of the cutoff, which give $\nu = 0.60$ for $m_{\text{min}} = 6000$, $\nu = 0.59$ for $m_{\text{min}} = 7000$ and 8000, and $\nu = 0.58$ for $m_{\text{min}} = 9000$. Hence we fix $\nu$ to its theoretical value and fit the correction-to-scaling exponent to all available data (setting a lower cutoff increases sensibly the errors, but the values are all compatible with the one given below). The results of this procedure (plotted in Fig. 5) bring to

$$\Delta_1 = 0.76(5). \quad (14)$$

This exponent, to our knowledge, has never been computed for $\theta$-point polymers, and its value in percolation has been the subject of long debate \cite{37}. Our result $\Delta_1 = 72/91 \approx 0.791$ obtained independently from calculations in the Potts model \cite{35} and in the $O(n)$ model \cite{39}. However, caution must be exercised; as already noted, corrections to scaling are a notoriously slippery ground, since terms with different exponents (and possibly different signs) can conspire to yield deceptive effective exponents. Further confirmations of this result, perhaps obtained with other techniques, are awaited.

Although rotational symmetry forces the distribution functions to be spherically symmetric, the instantaneous shape of polymers is not spherical on average. A measure of the deviations from spherical shape is given by the gyration tensor

$$Q_{\alpha\beta} = \frac{1}{2(n+1)^2} \sum_{i,j=0}^n \left( \gamma_{i,\alpha} - \gamma_{j,\alpha} \right) \left( \gamma_{i,\beta} - \gamma_{j,\beta} \right), \quad (15)$$

where $\gamma_{i,\alpha}$ is either $\text{Re}(\gamma_i)$ or $\text{Im}(\gamma_i)$ depending on the value of $\alpha = 0, 1$ (note that it implicitly depends on $n$). $Q_{\alpha\beta}$ is symmetric and positive definite; therefore it has two positive eigenvalues, $q_1$ and $q_2$, say such that $q_1 \geq q_2$. Two quantities characterizing the shape of the curves can be constructed from the eigenvalues:

$$r_n = \frac{q_1}{q_2} n, \quad A_n = \frac{ \left( q_1 - q_2 \right)^2 }{ \left( q_1 + q_2 \right)^2 } n. \quad (16)$$

called shape factor and asphericity respectively. For a spherically symmetric object, $r = 1$ and $A = 0$, while in the extreme case of a rod-like curve the asphericity is 1 and the shape factor diverges. These quantities are expected to have a finite limit for large $n$, and behave as $r_n \sim r_{\text{SLE}} - a/n^{\Delta_1}$ and $A_n \sim A_{\text{SLE}} - b/n^{\Delta_1}$ to leading order in $n$.

We analyzed the shape factor and the asphericity for the SLE chains, by performing independent simulations at $n = 1000, 2000, 3500, 5000, 6000, 7500, 8500, 10000, 15000, 20000$. We remind that the chains are generated in such a way as to have constant step lengths. Approximately 100,000 samples were produced for each $n$, except $n = 5000, 10000$ (200,000 samples) and $n = 15000, 20000$ (about 50,000 samples). Results are presented in Fig. 6. Unexpectedly, we find that the predicted scaling form is violated considerably for $n \lesssim 5000$. Therefore we fix a lower cutoff and fit the results only for $n > 5000$. From the fits we obtain

$$r_{\text{SLE}} = 4.38(3), \quad A_{\text{SLE}} = 0.373(1), \quad (17)$$

which shows that the curves are strongly elliptical. The estimated values of $\Delta_1$ are affected by huge errors ($\Delta_1 = 1.8 \pm 1.0$ and $2.4 \pm 1.6$, for $r$ and $A$ respectively) and are thus not comparable to theoretical predictions, but they are compatible with the more precise estimate obtained above from the end-to-end radius (repeating the fits by keeping $\Delta_1$ fixed to an extreme of its confidence interval yields $r_{\text{SLE}} = 4.33 \sim 4.42$ and $A_{\text{SLE}} = 0.370 \sim 0.377$). The asymptotic values are surprisingly close to those of ordinary $\theta$-point polymers in two dimensions; in fact, they

Figure 5: (Color online) The asymptotic behavior of the end-to-end radius for SLE chains is consistent with that of $\theta$-point polymers. The effective exponent $\nu_{\text{eff}}$ measured from simulations (red crosses) is plotted against the chain length $n$; the dotted blue line is the asymptotic value corresponding to the $\theta$-point prediction; the dashed black line includes the fitted corrections to scaling. The inset shows the end-to-end radius, from which the effective exponent is computed, together with its asymptotic expression.
are compatible within statistical significance. The best estimates for polymers, depicted by the shaded intervals in Fig. 6 are $r_θ = 4.46(6)$ and $A_θ = 0.3726(7)$ [28]. By comparison, self-avoiding walks in the plane and discrete SLE curves with constant step lengths at $κ = 8/3$ have $A_{SAW} ≈ 0.5134$ [10], while for true random walks $A_{RW} ≈ 0.3964$ [30]. We do not have an explanation for the coincidence of the asphericity measures. As already noted, the influence of the finite-chain effects and of the domain (the plane punctured by a disc) modifies the shape and the distribution functions (Fig. 4). The asphericity for SAW has already been shown to have a non-trivial dependence on the position of the internal point [10]. Therefore the similarity in the overall shape that we observe for $κ = 6$ and the $θ$ point is intriguing. Our result for the exponent $Δ_1$, instead, fits into the theoretical picture, and shows the power of the numerical method. As already noted, the SLE-based numerical strategy applied to the case $κ = 8/3$ [10] exposed, for self-avoiding paths, the presence of leading corrections to scaling with the exponent predicted by conformal field theory, $11/16$, which are surprisingly absent for square-lattice SAW [28] (where the leading corrections have exponent $3/2$). In this perspective, it would be interesting to calculate $Δ_1$ directly for $θ$-point ISAW, and also for their endless counterpart on the lattice, as was done in [9] for non-interacting SAW.

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