Quantum criticality in SU(3) and SU(4) anti-ferromagnets

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We study the quantum phase transition out of the Néel state in SU(3) and SU(4) generalizations of the Heisenberg anti-ferromagnet with sign problem free four spin coupling (so-called JQ model), by extensive quantum Monte Carlo simulations. We present evidence that the SU(3) and SU(4) order parameters and the SU(3) and SU(4) stiffness go to zero continuously without any evidence for a first order transition. However, we also find considerable deviations from simple scaling laws for the stiffness and discuss scenarios to explain the observed behavior.

I. INTRODUCTION

The theory of quantum phase transitions has become a centerpiece in the research of quantum physics of strongly coupled condensed matter. It has found applications in many branches of condensed matter physics, examples include: heavy fermion systems, high temperature superconductivity, quantum Hall effect, metal-insulator transition, ultra-cold atomic gases and frustrated magnets. Aside from its many applications to natural systems, the study of quantum phase transitions has also turned into a full fledged theoretical endeavor in its own right, which although very mature in certain aspects, is still in a stage of infancy when one considers the large number of different situations modern condensed matter systems allow for. Indeed, most of the well understood quantum critical points (continuous quantum phase transitions) can be simply related to a classical universality class in one higher dimension. On the other hand, one of the most exciting directions in the field is the study of quantum critical points which lead to new universalities that are not natural to think of in the classical context. Such new criticality may arise, for instance, from essentially quantum phenomena such as the presence of low-energy fermions or complex Berry phases that have no natural classical analogue.

An interesting theoretical proposal for a quantum critical point that does not have a naive classical analogue was put forward a few years ago. The physical system is a two-dimensional quantum anti-ferromagnet on a square lattice with S=1/2 spins which transitions into a paramagnet with a broken translational symmetry, a valence bond solid (VBS). In a set of compelling arguments, it was shown that a naive application of the classical theory which forbids a continuous transition could be invalidated by the presence of Berry phase terms, giving rise to a quantum critical point in a novel universality class. It is widely accepted, however, that a full non-perturbative understanding of the occurrence and properties of the new quantum critical point are beyond analytic reasoning and can only be established by unbiased numerical simulations. In a pioneering piece of work, it was shown that these questions could be addressed in a sign problem free microscopic JQ model using quantum Monte Carlo techniques. Since then a number of works have studied this SU(2) symmetric model. While all workers agree that the JQ model harbors a Néel state on one side and a VBS state on the other side, a clear picture of the the nature and precise scaling at the quantum phase transition still does not exist. The most comprehensive numerical study of the SU(2) JQ model however provides strong evidence in favor of a continuous transition, albeit with corrections to the naïve scaling hypothesis. Corrections to naïve scaling also appear in the study of the response of the JQ quantum critical point to single impurities. Another study for an unconventional quantum critical point in the spatially anisotropic bilinear bi-quadratic model has also found evidence for a continuous quantum critical point, though this model is less well studied. The only theory for this transition is also beyond a naive classical order parameter theory. Excluding the JQ model and the anisotropic bilinear biquadratic $S = 1$ model, there is no known candidate for a quantum critical point in a numerically accessible two dimensional microscopic quantum Hamiltonian that does not map simply onto a higher dimensional classical field theory (the most well studied example of a quantum critical point with a higher dimensional classical mapping is the bilayer anti-ferromagnet, the mapping has been studied thoroughly). Sorting out these issues is a very demanding one numerically, but is extremely important for the field of quantum criticality.

In this paper we present the results of extensive QMC simulations on a larger $N$ extension of the original SU($N = 2$) symmetric JQ model. The first study of these models reported $T = 0$ QMC results using the valence bond basis method for the SU(3) and SU(4) models and found results consistent with a continuous transition with conventional scaling for quantities associated with the VBS and Néel order parameters. In this work we approach this model with the powerful finite-T stochastic series expansion (SSE) method which allows access to finite-temperature, larger volumes and most importantly for our purposes here, gives access to the stiffness and susceptibility. We find that these quantities have large corrections to scaling, just like those found in recent studies of the SU(2) symmetric JQ model. This result does not contradict the findings of conventional scaling in Ref. since their results were focused on the order parameters. Indeed for the Binder ratio associated with the Néel order parameter we too find conventional scaling works very well (see Sec. ). The corrections we
find are restricted to the susceptibility and stiffness (defined below), which are susceptibilities of the generators of the SU(N) symmetry and which cannot be accessed in valence bond basis method so far, which was used in the previous study. The major new result of this paper is a study of the possible scaling forms for these quantities (i.e. the stiffness) in the SU(3) and SU(4) symmetric models. We also present a number of scenarios which could explain the observed numerical results. Larger-N extensions of the SU(2) physics may have applications to cold atom systems, but are important in their own right as they allow one to approach the analytically well-understood large-N limit. Other numerical work has studied the phase transition between Néel-VBS by varying the N in SU(N) continuously, without the introduction of a four spin Q term.

II. MODEL

We define our model on a square lattice, which can be conveniently separated into an A and B sub-lattice. We define N flavors of fermions, \( f_\alpha \), identically on each site \((1 \leq \alpha \leq N)\). An SU(N) rotation will rotate among these N flavors with the following convention:

\[
f_\alpha \rightarrow f'_\beta = U_{\alpha\beta} f_\beta
\]  

(1)

where * implies complex conjugation. Now to pick a spin representation in terms of the \( f_\alpha \) fermions, we fix the number of fermions on each site, \( f'_\alpha f_\alpha = 1 \) on the A sub-lattice and \( f'_\alpha f_\alpha = (N - 1) \) on the B sub-lattice. This fixes the size of the local Hilbert space to be simply N on each site, on both the A and on the B sub-lattice. We define these N states with the following sign conventions and indicate their transformations under SU(N) rotation,

\[
|\alpha\rangle_A = f_\alpha^\dagger |0\rangle \quad (|\alpha\rangle_A \rightarrow U_{\alpha\beta} |\beta\rangle_A)
\]  

(2)

\[
|\alpha\rangle_B = f_\alpha |F\rangle \quad (|\alpha\rangle_B \rightarrow U_{\alpha\beta}^\dagger |\beta\rangle_B)
\]  

(3)

where \(|0\rangle\) denotes the absence of any fermions and \(|F\rangle\) is a fully filled site. The transformation properties imply that the N, A sub-lattice states transform in the fundamental representation of SU(N) and the N, B sub-lattice states transform in the conjugate of the fundamental representation. The transformations also imply that the state \( \sum_\alpha |\alpha\rangle_A |\alpha\rangle_B \) transforms as an SU(N) singlet. This is the main reason we have chosen different representations on the A and B sub-lattices, i.e. to allow the formation of a two-site singlet, following previous work on anti-ferromagnets.

To construct a spin model, we write down SU(N) invariant four fermion interactions that maintain the number of fermions on each site; Explicitly, there are two such terms \( f_\alpha^\dagger f_\alpha f^\dagger_j f_j \) and \( f_\alpha^\dagger f_\alpha f_j f^\dagger_j f_\alpha^\dagger \); the first one is just the identity operator in the projected space, while the second term which we call, \( P_{ij} \) is a projector onto an \( ij \) singlet (\( i \) is on A sub-lattice and \( j \) is on B sub-lattice).

Note that the matrix elements of \( P_{ij} \) are very simple,

\[
\langle \alpha_1 \beta_2 | P_{ij} | \alpha_2 \beta_2 \rangle = \delta_{\alpha_1 \alpha_2} \delta_{\beta_1 \beta_2}
\]  

(4)

Since they are always positive when non-zero, we can use these operators to construct sign problem free models. In particular the model defined by \( H_J = -\frac{J}{N} \), is the familiar Heisenberg model up to a constant, after identifying \( \alpha = 1 \) with up(down) and \( \alpha = 2 \) with down(up) on the A(B) sub-lattice. We define the SU(N) “JQ” model with the following conventions.

\[
H_{JQ} = -\frac{J}{N} \sum_{ij} P_{ij} - \frac{Q}{N^2} \sum_{ijkl} P_{ij} P_{kl}
\]  

(5)

where the first sum is taken over all nearest neighbor bonds on the square lattice and the second term is taken over all elementary plaquettes of the square lattice with \( ij \) and \( kl \) being nearest neighbor bonds. Note that so defined, \( H_{JQ} \) explicitly has no sign problem: all off-diagonal matrix elements are explicitly negative.

We now turn to the observables of interest in our study here. There are \( N^2 - 1 \) traceless Hermitian matrices, \( X^a_{\alpha\beta} \) which generate the SU(N) algebra. Of these, \( N - 1 \) can be chosen diagonal (so-called Cartan subalgebra). We choose them so that they satisfy: \( \text{Tr}[X^a X^b] = \delta_{ab}/2 \). We can then work out formulae for the “uniform magnetization” and “staggered magnetization” for the Cartan generators in terms of the operator \( n_\alpha \) which measures which of the \( |\alpha\rangle \) states is occupied on each site (\( n_\alpha = f_\alpha^\dagger f_\alpha \) on the A sub-lattice and \( n_\alpha = 1 - f_\alpha^\dagger f_\alpha \) on the B sub-lattice).

\[
M_a^a = \sum_{r_\in A} X^a_{\alpha\alpha} n_\alpha - \sum_{r_\in B} X^a_{\alpha\alpha} n_\alpha
\]  

(6)

\[
M_a^b = \sum_{r_\in A} X^a_{\alpha\alpha} n_\alpha + \sum_{r_\in B} X^a_{\alpha\alpha} n_\alpha
\]  

(7)

Note that the sign in the middle is opposite from what you would have expected for the familiar \( S = 1/2 \) case, because of the way our Hilbert space is defined on the A and B sub-lattices. The specific choices for the Cartan generators is detailed in Appendix A. An outline of the algorithm and method is given in Appendix A.

III. BINDER RATIO

The first quantity which we study is the so-called Binder ratio. It is defined as,

\[
B = 1 - \frac{\langle M_a^4 \rangle}{3\langle M_a^2 \rangle^2}
\]  

(8)

In addition to the Monte-Carlo averaging, the \( \langle \ldots \rangle \) imply averaging over the \( N - 1 \) Cartan generators. It is well known that the Binder ratio has a scaling dimension of zero and hence can be written in the following simple scaling form close to a quantum critical point,

\[
B = \mathbb{E}(\frac{L^{\nu}}{c}, gL^{1/\nu})
\]  

(9)
FIG. 1: A zoom-in of the crossing of the binder ratio, $B$, for SU(3) and SU(4) symmetric JQ models. Error bars were determined by bootstrapping the data. The solid lines are polynomial fits on the data and are plotted to guide the eye. The crossing point converges well for the larger system sizes. For $L = 64, 96, 128, 192$ the data crosses nicely within the estimated error bars, allowing very accurate brackets for the quantum critical points. $(J/Q)^2_\text{SU(3)} = (1.9905, 1.9920)$ and $(J/Q)^2_\text{SU(4)} = (11.235, 11.255)$.

This means that if we fix the parameter $LT$ (see Appendix C), we assume everywhere in this work that $z = 1$, that $B$ should become volume independent when $g = 0$ in the scaling limit (i.e. for large $L$). The simplest way to find whether this is true is to plot $B(g)$ for different values of $L$ and look for a crossing of the various curves. Such a plot for both the SU(3) and SU(4) JQ model is shown in Fig. 1 We have collected data on system sizes ranging from $L = 16 - 192$. While the data is accurate enough to detect corrections to scaling for the smaller sizes the crossing converges very well as the system size is increased and the four biggest system sizes cross nicely within our estimated error bars. From this crossing we have an accurate estimate of the value of the critical coupling (see caption of Fig. 1), which is consistent with the values quoted in Ref. [19] thanks to the larger volumes simulated here, our brackets are about a factor of 2 times more accurate for both SU(3) and SU(4)].

IV. STIFFNESS

Configurations of our SU($N$) model in the SSE method (see Appendix A) can be thought of as a set of $N$ colors of non-intersecting closed loops. We can measure the winding number of each of the $N$ colors of loops both in space and in time and associate a spatial and temporal current for each configuration, with each of the $N - 1$

diagonal generators.

\begin{align}
    j_x^a &= X_{\alpha}^a W^\alpha_x \\
    j^\tau &= X_{\alpha}^a W^\alpha_\tau
\end{align}

The fluctuations of these quantities averaged over MC sampling and the $N - 1$ diagonal generators $\langle W^2 \rangle \equiv \langle (j_x^a)^2 \rangle$ and $\langle W^2 \rangle \equiv \langle (j^\tau)^2 \rangle$, are of great interest, since they too are expected to have no scaling dimension just like the Binder ratio, $B$, discussed earlier, because they are susceptibilities of the generators of a conservation law (SU($N$) symmetry). Hence, for the same reasons discussed in the case of $B$, they are expected to have a crossing point. Before turning to a numerical study of their crossing, we note that by linear response theory, the fluctuations in spatial winding number are simply related to the familiar spin stiffness ($\rho_s = \langle W^2 \rangle / \beta$) and the fluctuation in the temporal winding number are related to the spin susceptibility ($\chi_u = \langle W^2 \rangle / \beta L^2$), we shall refer to these interchangeably.

Fig. 2 shows $\langle W^2 \rangle (g)$ for different $L$, for both the SU(3) and SU(4) model. While the data does cross, it does so very roughly. There are clearly large corrections to scaling behavior. Are these just corrections to scaling or is the leading scaling behavior itself affected? To make further progress in understanding the deviations from scaling, we study the crossing point of $\langle W^2 \rangle$ for the sizes $L$ and $2L$ in Fig. 2. The crossing points have an $x$-intercept (the coupling $J/Q$) and a $y$-intercept (the value of $\langle W^2 \rangle$). We study these intercepts as a function of $L$ with the hope of being able to extrapolate the large-$L$ behaviors. In order to estimate an error bar for these crossing points, we carry out a somewhat labori-
ous bootstrap method, in which we generate synthetic data sets of the same length as our data (drawn by randomly sampling our own data), carry out least squares fit to this data (with polynomials of order two), and then compute the crossing point in each simulated data set. The RMS of the extracted crossing point data is then a reasonable estimate of the error in determining the $x$- and $y$-intercepts of the crossing point. Fig. 3 shows the $J/Q$ value for the crossing points as a function of $1/L$. The data clearly converges for large $L$ to a finite value. It is re-assuring that this value is completely consistent with the brackets for the critical point extracted from the Binder ratio data (shown as thick lines on the $y$-axis of Fig. 3). Next, we turn to the $y$-intercept of the crossings. Here things are much harder to interpret because the data does not saturate even for the largest system sizes studied here. Of course, one could always argue that this is due to a finite size effect, but given the large volumes simulated here, and the absence of any sign of saturation in the $y$-intercept, we shall assume that $\langle W_2^2 \rangle$ diverges as the volume increases. We would like to note here that this by itself does not indicate a first order transition. At a first order transition $(\langle W_2^2 \rangle)$ should grow linearly (because the stiffness itself remains finite instead of going to zero). We see in the insets of Fig. 3 that there is no evidence for a linear divergence. We find that for our data on the SU(3) and SU(4) models, a weak power law divergence seems to fit better than the logarithmic divergence found for SU(2) [14], see Figs. 4 and 5.

V. DISCUSSION

It is interesting to understand the theoretical reasoning behind the numerical analysis presented here. While we cannot offer an unambiguous theoretical interpretation, we make some inferences on what could be taking place. First we summarize our conclusions from what we have observed,

- There is essentially no difference in the qualitative behavior for the various quantities under study from SU(2) for both SU(3) and SU(4). Quantitative differences in quantities such as the critical exponents must clearly exist as the symmetries change.

- The Binder ratio crossing seems to follow the naive scaling expectation of a quantum critical point with one relevant direction and all other directions irrelevant. There are corrections to scaling but they are of the conventional type, i.e., they do not appear to affect the leading scaling behavior and vanish for large system sizes.

- The stiffness (and susceptibility) have large corrections to scaling. The crossing point converges on the $x$-axis to the same critical coupling deduced from the Binder ratio, whereas the value of the $L_{\rho_0}$ itself ($y$-axis) has a sizable volume dependence. If we assume the observed behavior in Fig. 2 is the leading scaling behavior, we conclude that the winding number fluctuations diverges sub-linearly. The fact that the divergence is sub-linear is very crucial, since it implies that the stiffness (which is the winding number fluctuation divided by $L$) itself goes to zero as one approaches the critical point, albeit not as $1/L$ like one would have expected from a naive scaling analysis. It is difficult numerically to distinguish between a log or a very weak power law divergence, as we possibly have here, since the $y$-axes in Fig. 4 do not change very significantly over the entire range of sizes studied.

Perhaps, the simplest explanation for the stiffness measurement is that the volume dependence of the stiffness curve is a finite-size effect and that it would saturate as the system size is made even larger than that studied here. This would be consistent with a conventional scaling hypothesis for the critical points studied here. While we cannot rule this possibility out, we believe it is unlikely since the crossing in the Binder ratio seems to have converged well for the system sizes under study. If this explanation is to be made viable, one would need a theoretical explanation for why the finite-size corrections to

\[ FIG. 3: \text{The} \ x\text{-intercept} (J/Q) \text{of the crossing of} \ L \text{and} 2L \text{for the stiffness shown in Fig. 2 plotted versus} \ 1/L. \text{The errors bars are estimated by a bootstrap method detailed in the text. The error bars get smaller for large systems because these curves are steeper; the} x\text{-intercept of the crossing can hence be determined more precisely. At} 1/L = 0 \text{the thick red line shows the bracket for the critical point from the analysis of the Binder ratio data (Fig. 1). Clearly the} x\text{-intercept of the crossing converges to a finite value, and this value is completely consistent with the Binder ratio crossing. The dashed line is a non-linear fit to the form} g - A/L^a, \text{with} g = 1.992, A = 3.20, a = 1.47 \text{for SU(3) and} g = 11.255, A = 51.3, a = 1.48. \text{A motivation for this form is given in Appendix D.} \]
the stiffness are as large as they are.

If we assume that the observed behavior is indeed the asymptotic behavior, we are faced with another vexing questions: What could result in this unusual behavior for the stiffness? This behavior must signal the breakdown of one of the standard scaling assumptions made. One scenario which would result in the divergence of $L \rho_s$ is if the scaling function depended non-analytically on an irrelevant operator. For instance, imagine that $L \rho_s = f(g L^{1/\nu}, \frac{x}{L})$ with $\omega > 0$ and hence $\phi_\omega$ formally irrelevant. Normally one assumes the scaling function is analytic in its arguments and can hence set the second argument to zero. If it was not analytic and depended for instance like $f(x, y) \approx \frac{1}{y}$ for small $y$, there would be a power law divergence in $L \rho_s$ as observed here. One important consequence of this scenario is that a power law divergence is more natural than a log divergence, since a log divergence (within this scenario) would require a singular dependance of the scaling function and an exactly marginal operator in addition. Of course this scenario is completely speculative and if it is indeed true, we leave it for future work to understand what this irrelevant operator is and why it results in singular scaling functions for the stiffness.

An interesting discussion is how the correction evolves from the SU(2), SU(3) to SU(4) critical points. Since the first theoretical work about such critical points, it has been assumed that the properties of the critical point depend smoothly on $N$. Our findings are not inconsistent with that assumption. While it is numerically difficult to distinguish between a weak power law (as we might have here) and a logarithm, there are important theoretical distinctions that arise from such behavior. In particular, if the log behavior found for SU(3) results from a marginal operator, it is natural to assume that the scaling dimension of this operator should depend on $N$. This would make it hard to explain a log divergence in the SU(3) or SU(4) critical points, even though such behavior is not inconsistent with our numerical data. It could also be possible that in the SU(2) case a weak power law divergence of the same form as proposed here for SU(3) and SU(4) is present. A complete theoretical scenario is required to address these interesting possibilities.

VI. ACKNOWLEDGMENTS

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FIG. 6: An identical analysis as that leading up to Fig. 5 but now for the fluctuations of the temporal winding number. The power law “fit” does not work quite as well here. Again the “fit” here is carried out by fitting the power law form $Ax^B$ to the two largest system sizes. The exponents we find here are $B_3 = 0.18(2)$ and $B_4 = 0.17(2)$, where the errors are determined roughly by fitting different sets of the data. Re-assuringly, they are in agreement with the values found for the stiffness. See also Fig. 7 for a comparison between the susceptibility and stiffness.

Motrunich, and T. Senthil. All the simulation reported here were carried out on the BCX cluster at the Center for Computational Sciences at the University of Kentucky.

Appendix A: Method and algorithm

The method employed here is the stochastic series expansion, which is well documented in the literature. In the way we have defined the Hilbert space, one can think of the configuration space as a set of loops (the loops travel vertically up in time on the A sub-lattice and vertically down in time on the B sub-lattice) with a color assignment from one of $N$ colors. For the updates, we have generalized the “deterministic” method of Ref. 20, Sec. II D, to both the four site Q term and SU(N), this is possible since as noted in Sec. II because of the simple form of the matrix elements we are guaranteed that all configurations which appear at a given order of $J$ and of $Q$, appear with the same weight. In this method there are two kinds of updates (1) [loop update] the $N$ colors of a loop can be randomly assigned a new color and (2) [diagonal update] swap of identity operators with diagonal operators.

Appendix B: Choice of Cartan algebra

In this appendix we list our choices for the $N-1$ diagonal generators for $N = 3, 4$. There are different conventions with which these can be chosen. For SU(2) there is only one diagonal generator and the natural choice is:

$$
\begin{pmatrix}
1/2 & 0 \\
0 & -1/2
\end{pmatrix}
$$

for SU(3) we used:

$$
\begin{pmatrix}
1/2 & 0 & 0 \\
0 & -1/2 & 0 \\
0 & 0 & 0
\end{pmatrix}
$$

Finally for SU(4) we used:

$$
\begin{pmatrix}
1/(2\sqrt{3}) & 0 & 0 & 0 \\
0 & 1/(2\sqrt{3}) & 0 & 0 \\
0 & 0 & 1/(2\sqrt{2}) & 0 \\
0 & 0 & 0 & -1/(2\sqrt{2})
\end{pmatrix}
$$

$$
\begin{pmatrix}
-1/(2\sqrt{2}) & 0 & 0 & 0 \\
0 & 1/(2\sqrt{2}) & 0 & 0 \\
0 & 0 & 1/(2\sqrt{2}) & 0 \\
0 & 0 & 0 & -1/(2\sqrt{2})
\end{pmatrix}
$$

$$
\begin{pmatrix}
-1/(2\sqrt{2}) & 0 & 0 & 0 \\
0 & -1/(2\sqrt{2}) & 0 & 0 \\
0 & 0 & 1/(2\sqrt{2}) & 0 \\
0 & 0 & 0 & 1/(2\sqrt{2})
\end{pmatrix}
$$

All these choices are made so that they satisfy the standard normalization, $Tr[X^aX^b] = \delta_{ab}/2$.

Appendix C: Choice of Cartan algebra

Assuming $z = 1$ scaling, a choice has to be made for the ratio of $LT/c$. Of course any fixed ratio will work for the scaling properties, but in order to best use the efforts of our simulations we would like to work as close to the cubic limit as possible $LT/c = 1$. The problem is that we do not know what $c$ is in terms of our couplings $J$ and $Q$, a priori. To circumvent this problem, we pick the temperature so that the fluctuations of the winding number in space (the re-scaled spin stiffness) and the fluctuations in the winding number in time (the rescaled spin susceptibility) are of the same order of magnitude. Indeed
we would expect them to be identical at the isotropic point. This is achieved most simply by choosing a coupling close to the critical point and the choosing some appropriate system size (small is good enough, so that the correlation length is larger than the system size) and then adjusting the temperature till the fluctuations of the temporal and spatial winding number become approximately equal. We found that working in units with $Q = 1$ that the approximate equality of the temporal and spatial winding numbers was achieved for $L/\beta = 4$ for SU(3) and for $L/\beta = 10$ for the SU(4) case, we hence used these values for the simulations reported in this manuscript.

### Appendix D: Crossing point

In this appendix we study how the crossing analysis is affected by a multiplicative correction to the standard scaling function.

Imagine we had a scaling function that had the form, $L^\alpha f(gL^\beta)$. Now let's ask for what value of $g_x$ it crosses. Assuming the function $f$ is analytic in its argument we find close to the critical point, $g_x \approx f(0)/L^\alpha f'(0)$. So, the value of the critical coupling does go to zero (the true critical point) for larger and larger $L$, but it already receives corrections without any corrections from irrelevant operators. The leading behavior of the quantity at the crossing point, described by such a scaling function is simply, $L^\alpha f(0)$.

In order to test the functional form of $g_x$, we have carried out a non-linear fit to the data on the stiffness as detailed in the caption of Fig. 3. We find $\nu \approx 0.65(5)$ for both SU(3) and SU(4) which is consistent with the value of $\nu$ reported in a previous study (the error quoted here for $\nu$ has been determined very roughly by attempting a number of non-linear fits dropping different sets of data points). Interestingly the $\nu$ in the previous work was extracted completely from the data on the order parameters. This gives some credibility to the idea that there is a multiplicative correction to the naive scaling for the stiffness. The analysis is very similar for $\ln(L)f(gL^\alpha)$, where again $g_x \approx \frac{f(0)}{\ln(L)L^\alpha f'(0)}$ for large $L$ and the function would be described by, $\ln(L)f(0)$ at leading behavior. Our $g_x$ data would fit this model reasonably too, since the multiplication of a log affects the numerical values very weakly.

### Appendix E: Susceptibility

In this appendix we study the fluctuations of the temporal winding number (the re-scaled susceptibility). We carry out an identical analysis to that presented in the body of the text for $\langle W_x^2 \rangle$.

The susceptibility has an identical crossing behavior to that of the stiffness shown in Fig. 2 there is a drift in the crossing point in both the $x-$ and $y-$intercepts. We have carried out an identical analysis of the crossing of $L$ and $2L$ and we find that the $x-$intercepts (shown in Fig. 3 for the stiffness), do not depend very much on whether we look at the crossing of the susceptibility or the stiffness. The $y-$intercept does however, so we have made a separate plot for this quantity in Fig. 4. We find that unlike in the case of stiffness there are finite-size correction to the power law model. Yet the power of the fitted power law for the largest system sizes is in agreement with our findings for the susceptibility, as detailed in the inset.

Finally, as an independent check that these two quantities diverge the same way as we approach the critical point, we have plotted the ratio of these two quantities at the crossing point value determined by the susceptibility crossing. This is shown in Fig. 7. As expected we find that the ratio of the quantities clearly goes to a constant at large volume, providing clear evidence that the two quantities diverge the same way, as one would expect for the emergent $z = 1$ scaling (we use the word emergent here, since the microscopic model is clearly not invariant under space-time rotations!).

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