Finite-temperature phase transitions in $S = \frac{1}{2}$ three-dimensional Heisenberg magnets from high-temperature series expansions

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Many frustrated spin models on three-dimensional (3D) lattices are currently being investigated, both experimentally and theoretically, and develop new types of long-range orders in their respective phase diagrams. They present finite-temperature phase transitions, most likely in the Heisenberg 3D universality class. However, the combination between the 3D character and frustration makes them hard to study. We present here several methods derived from high-temperature series expansions (HTSEs), which give exact coefficients directly in the thermodynamic limit up to a certain order; for several 3D lattices, supplementary orders than in previous literature are reported for the HTSEs. We introduce an interpolation method able to describe thermodynamic quantities at $T > T_c$, which we use here to reconstruct the magnetic susceptibility and the specific heat and to extract universal and nonuniversal quantities (for example, critical exponents, temperature, energy, entropy, and other parameters related to the phase transition). While the susceptibility associated with the order parameter is not usually known for more exotic long-range orders, the specific heat is indicative of a phase transition for any kind of symmetry breaking. We present examples of applications on ferromagnetic and antiferromagnetic models on various 3D lattices and benchmark our results whenever possible.

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I. INTRODUCTION

The quantum Heisenberg model was first introduced to explain why certain compounds developed a spontaneous magnetization when cooled below a given temperature (called the critical temperature), even in the absence of an applied magnetic field. Its success in explaining these so-called ferromagnets placed the model at a central spot in the study of quantum magnetism. Since then, a wide variety of threedimensional (3D) compounds have been investigated, whose interactions can be mostly described by ferro- or antiferromagnetic Heisenberg interactions. More often than not, these systems present finite-temperature phase transition to long-range magnetically ordered phases. Even in 2D systems, where the Mermin-Wagner theorem [1] precludes the existence of magnetic order at finite temperatures, materials often present finite-temperature phase transitions due to remaining weak 3D correlations [2–5]. Many of these phase transitions belong to the 3D Heisenberg universality class, defined by the symmetry breaking $SU(2) \rightarrow U(1)$.

Phase transitions are characterized by the critical temperature T_c and exponents of the singularities present in the thermodynamic functions. Obtaining them is thus an important task, and to do so many different methods have been developed. In particular, in the ferromagnetic case, hightemperature series expansions (HTSEs) methods such as the Dlog Padé or ratio methods are known to obtain numerically accurate critical temperatures for simple lattices such as the simple cubic (*sc*), body-centered cubic (*bcc*), and face-centered cubic (*fcc*) [6–10]. More recently the list was expanded with the pyrochlore [11], the diamond [10,12], and the semi-simple cubic (*ssc*) [13] lattices. However, for the critical exponents, the results are not so precise.

For example, the critical exponent γ of the magnetic susceptibility $\chi(T)$ has been calculated using field theory's renormalization group on the N-vector model (that is in the same universality class for N = 3), yielding $\gamma = 1.3895(50)$ [14,15]. On the other hand, HTSEs calculations in the quantum spin- $\frac{1}{2}$ case show higher values, $\gamma = 1.42(1)$ for the ferro- and $\gamma = 1.43(1)$ for antiferromagnetic cases [7–9]. Another example of a critical exponent is the less studied α from the specific heat $c_v(T)$. This exponent is negative, which implies a nondivergent singularity. Instead, $c_v(T)$ presents a cusp-like behavior that reaches a maximum value with an infinite slope. To the best of our knowledge, the standard HTSEs Dlog Padé and ratio methods have never been used on c_v in the literature. However, indirect HTSEs calculations through scaling relations give α between -0.125 and -0.200(15)[6,10,16]. On the other hand, the field theory result is $\alpha =$ -0.122(10) [15], showing a larger discrepancy than in the case of γ . On the experimental side, there are empirical fits on specific heat measurements that give negative values of α down to -0.3, while most of them lie around the field theory result [17,18].

Finally, it would be desirable to have reliable results not only on the critical quantities (critical temperature T_c and exponents) but on the thermodynamic functions at all temperatures $[c_v(T), \chi(T)]$. In this sense, the quantum Monte Carlo (QMC) calculations obtain reliable results, but only on finite lattices, and only in the absence of frustration due to the sign problem. Methods based on exact diagonalization and tensor network algorithms can be used on frustrated systems, but only in dimensions 1 and 2 [19-22]. Other methods work directly in the thermodynamic limit like the rotationally invariant Green's function method, which obtains qualitatively good results in 3D [23,24]. The pseudo-Majorana functional renormalization group provides quantitatively good results down to moderate temperatures but becomes uncontrolled at low temperatures [25–27]. Finally, the HTSEs are quasi-exact at high temperatures but fail close to the transition temperature, even when using Padé approximants. An interpolation scheme of HTSEs solving this was proposed in our previous article in cases where c_v presents a logarithmic divergence, such as in the 2D-Ising or 2D-XXZ models [28], resulting both in an evaluation of critical quantities and of $c_{\nu}(T)$ for temperatures from infinite down to T_c .

In this article, we first revisit the HTSEs Dlog Padé method for Heisenberg ferromagnets on several 3D lattices such as the *fcc*, *bcc*, *sc*, pyrochlore, and *ssc* lattices. For most, we calculated higher orders in the HTSEs than in previous articles, using an optimized algorithm. Also, we extend the Dlog Padé method to obtain quantities such as the critical energy e_c , the c_v critical exponent α , and nonuniversal quantities A and $B [c_v(\beta) \sim B - A(\beta_c - \beta)^{-\alpha}]$. Finally, we extend the previously mentioned interpolation method [28] to cases where c_v presents a cusp-like behavior with a negative exponent $\alpha > -1$ and to cases where χ presents a divergent singularity with positive exponent $\gamma > 0$. For some 3D lattices, we can extrapolate c_v and χ at all temperatures down to T_c .

The remaining of the article is organized in the following way. In Sec. II, we present the HTSEs methods to study finite-temperature phase transitions, both the Dlog Padé and our interpolation method. In Sec. III we show our results, first for the Dlog Padé, and next for the interpolation methods. Finally, conclusions and perspectives are given in Sec. IV.

II. MODEL AND METHODS

The Heisenberg model is defined as

$$\mathcal{H} = J \sum_{\langle ij \rangle} \mathbf{S}_i \cdot \mathbf{S}_j, \tag{1}$$

where *J* is the exchange interaction, the sum $\langle ij \rangle$ runs over nearest neighbors on a 3D lattice, and **S**_i are the quantum spin- $\frac{1}{2}$ operators. The classical approximation consists in replacing the operators **S**_i by 3D vectors. In the ferromagnetic case (*J* < 0), the quantum ground-state energy per site e_0 is exactly the same as the classical one, namely,

$$e_0 = -\frac{Z}{2}S^2 \tag{2}$$

where Z is the coordination number of the lattice. Even though e_0 does not change when taking into account quantum fluctuations, the critical temperature T_c does [8]. On the other hand, in the antiferromagnetic case (J > 0) on a bipartite lattice, e_0 and T_c are the same as in the ferromagnetic case at the classical

limit, but they both change in the quantum model and e_0 is no longer known exactly.

To study these kinds of finite temperature phase transitions we use HTSEs. HTSEs allow to perform a series expansion of certain thermodynamic functions around $\beta = 0$, where β is the inverse temperature ($\beta = 1/T$). Two important functions are the free energy per site f and the ferromagnetic zero-field susceptibility per site χ [7]. Their HTSEs are written as

$$\beta f = -\ln 2 - \frac{1}{n_u} \sum_{i=1}^n \frac{a_i}{4^i i!} K^i + O(K^{n+1}), \qquad (3a)$$

$$\overline{\chi} = T\chi = \frac{1}{4} + \frac{1}{2n_u}\sum_{i=1}^m \frac{b_i}{4^i i!}K^i + O(K^{m+1}),$$
 (3b)

where a_i and b_i are integers, $K = \beta J$, and n_u is the number of spins in the unit cell. These HTSEs are typically known up to orders 13 to 15 for 3D lattices (see Table I for the order depending on the lattice).

The thermodynamic functions present singularities at the critical temperature. However, several methods can be used to extract information about the critical point from the first coefficients of the series. We present in Sec. II A the most commonly used: the Dlog Padé method, and pursue in Sec. II B with the description of a new interpolation method.

A. Dlog Padé method

We assume a thermodynamic function f(x) that has a power-law singularity at x_c , of type

$$f^{s}(x) = A(x_{c} - x)^{-\theta},$$
 (4)

such that $f(x) - f^s(x)$ is analytic from x = 0 to some $x > x_c$. x_c is the critical point and θ is the critical exponent. Then, the critical point and exponent can be obtained from the Dlog Padé method: the logarithmic derivative

$$D\ln f^{s}(x) = \frac{f^{s'}(x)}{f^{s}(x)} = \frac{\theta}{x_{c} - x}$$
 (5)

has a simple pole given by x_c , whose residue is the critical exponent θ . In practice, the critical point and exponent are determined from the poles and residues of the Padé approximants of the HTSE of $D \ln f(x)$.

This method has been used with $f(x) = \overline{\chi}(\beta)$ to obtain results for β_c and the critical exponent γ on most of the typical 3D lattices [7,8,10]. The Dlog Padé method presents a fast convergence of β_c with the HTSE order, giving several significant digits. Notably, to the best of our knowledge, this method has never been used with other thermodynamic functions such as c_v to determine α , or to obtain the critical values of the energy e_c and of the entropy s_c , what is now done in Secs. III B and III C.

B. Interpolation method for cusp singularities

Now we propose an alternative method to extract information on the critical point, using the specific heat. This is an extension from our previously introduced interpolation method for the case of logarithmic singularities [28]. In the present case (3D Heisenberg universality class), the singular behavior of the specific heat is expressed as

$$c_v^s(\beta) = -A(\beta_c - \beta)^{-\alpha} \tag{6}$$

where A is positive. Also, $-1 < \alpha < 0$ so that there is no divergence at the critical point. Instead, c_v reaches a maximum value B with an infinite slope (from higher temperatures). Other confluent terms, leading to more accurate results in the classical case [16], are less important in the quantum case [7].

We build a regular function $R(\beta)$ by removing the singular behavior from the specific heat. We explore two different ways of doing this. The first one, called the *interpolation method 1* (IM1), is analogous to that of Ref. [28],

$$R(\beta) = c_v(\beta) - c_v^s(\beta). \tag{7}$$

With this definition, the value *B* of c_v at the singularity is $R(\beta_c)$. The c_v -HTSE coefficients are calculated for a specific model up to an order *n*, and the series of c_v^s are known at all orders supposing that β_c , *A*, and α are known. Thus, the *R*-HTSE are obtained at order *n*, whose coefficients depend on β_c , *A*, and α . Compared to the case with a logarithmic divergency [28], the parameter space has one more dimension.

The other alternative to build the regular function R, called *interpolation method* 2 (IM2) is

$$R(\beta) = \frac{1}{A} \frac{c_v^s(\beta)}{c_v(\beta) - B}.$$
(8)

Again, the *R*-HTSE can be obtained up to order *n* and this time, it depends on β_c , *B* (instead of *A*), and α . Defined this way, $R(\beta_c) = 1/A$. It is worth mentioning that other similar methods could be developed with other regular functions *R*. But the two proposed here are sufficiently different so if both methods give similar results, we consider the results as trustworthy.

The idea behind these kinds of methods is that the parameters to obtain $R(\beta)$ have to be well chosen for R to be a truly regular function. This means that the singularity has to be canceled exactly. When this is done, the Padé approximants of $R(\beta)$ will coincide down to the critical temperature (and a little further below). The quality of a given set of parameters { β_c , A, α } (for IM1) or { β_c , B, α } (for IM2) is measured by the quality function already introduced in Ref. [28],

$$Q^{2} = \frac{2}{(n-1)n} \sum_{i=1}^{N_{\mathcal{P}}} \sum_{j=1}^{i-1} M_{\epsilon} \left(\frac{\mathcal{P}_{i}(\beta_{m}) - \mathcal{P}_{j}(\beta_{m})}{\overline{F}(\beta_{c})} \right) \quad (9)$$

where $N_{\mathcal{P}}$ is the number of Padé approximants \mathcal{P}_i without singularities in the range $[0, \beta_m]$, β_m is chosen larger that β_c to check the regular character of R beyond the critical point. We take $\beta_m = (1 + \delta)\beta_c$ with $\delta = 0.05$. $M_{\epsilon}(x)$ is a smooth function whose value is ≤ 1 when $x \ll \epsilon$, and ≥ 0 when $x \gg \epsilon$. We use $M_{\epsilon}(x) = 1/[1 + (x/\epsilon)^8]$ with $\epsilon = 0.005$. Finally, $\overline{F}(\beta_c) = \frac{1}{2}[\mathcal{P}_i(\beta_c) + \mathcal{P}_j(\beta_c)]$ is the average of the two Padé approximants at β_c . This Q function represents roughly the proportion of coinciding Padé approximants down to the critical temperature. Parameters with Q > 0.5 are considered as good.

Once a high-quality set of parameters is found, $c_v(\beta)$ can be reconstructed by replacing the regular function with any of TABLE I. The order *n* of the HTSEs used for βf and $\overline{\chi}$ in this article (we use the same order for both quantities), compared to orders from other articles. The lattices are *fcc* (face-centered cubic), *bcc* (bond-centered cubic), *sc* (simple cubic), *ssc* (semi-simple cubic), and the pyrochlore lattice. *sc-ssc* is a model interpolating between *sc* and *ssc*, defined in the main text. The new orders are provided in Appendix.

Lattice	This article	$n_{eta f}$	$n_{\overline{\chi}}$
fcc	13	12 [7]	14 [10]
bcc	15	14 [7]	14 [7]
SC	17	14 [7]	14 [7]
SSC	20		14 [13]
pyrochlore	16	13 [<mark>30</mark>]	12 [30]
<i>sc-ssc</i> , Eq. (18)	13		

its coinciding Padé approximants \mathcal{P}_i ,

$$c_{v}(\beta) = \mathcal{P}_{i}(\beta) - A(\beta_{c} - \beta)^{-\alpha} \quad \text{for IM1}, \qquad (10a)$$

$$c_v(\beta) = B - \frac{(\beta_c - \beta)^{-\alpha}}{\mathcal{P}_i(\beta)} \qquad \text{for IM2.} \qquad (10b)$$

C. Interpolation method for divergent singularities

The two methods presented in the previous subsection can be extended to quantities with divergent singularities, such as the magnetic susceptibility, whose singular part writes

$$\overline{\chi}^{s}(\beta) = C(\beta_{c} - \beta)^{-\gamma}$$
(11)

where *C* and γ are positive. From this point, the two methods IM1 and IM2 can be applied as in the previous subsection with the simplification that no constant term has to be taken into account (the term *B* of the previous section can be discarded as the divergency dominates it). This leads to an important difference between the extensions of IM1 and IM2 to divergent singularities. For IM1, *C* has to be taken into account and the parameter space consists in { β_c , *C*, γ }. But for IM2,

$$R(\beta) = \frac{1}{C} \frac{\overline{\chi}^{s}(\beta)}{\overline{\chi}(\beta)} = \frac{(\beta_{c} - \beta)^{-\gamma}}{\overline{\chi}(\beta)},$$
 (12)

the parameter space is reduced to $\{\beta_c, \gamma\}$. Thus, we will only use IM2 to interpolate χ in the following. From this regular function, the rest of the method is the same as described in the previous subsection, and the susceptibility can be reconstructed from any of its coinciding Padé approximants,

$$\chi(\beta) = \beta \frac{(\beta_c - \beta)^{-\gamma}}{\mathcal{P}_i(\beta)}.$$
(13)

III. RESULTS

We have numerically but exactly calculated the HTSEs of $\beta f(\beta)$ and $\overline{\chi}(\beta)$ for several 3D lattices: the *fcc*, *bcc*, *sc*, *ssc*, and pyrochlore lattices. The maximum order *n* depends on the lattice according to Table I, where we get the same order for both $\beta f(\beta)$ and $\overline{\chi}(\beta)$. Using an improved algorithm [29], we are able to calculate several orders more than previous studies for different lattices [7,10,13,30]. The new terms in the HTSEs are provided in Appendix.



FIG. 1. Density of poles as defined in Eq. (14) from the Dlog Padé method on $\overline{\chi}(\beta)$ for the ferromagnetic cases on the *fcc*, *bcc*, *sc*, *ssc*, and pyrochlore lattices as a function of $\beta - \beta_c$. For each curve, the poles from the four highest orders are taken.

In what follows, we study mostly the ferromagnetic Heisenberg model [J = -1 in Eq. (1)] on said lattices. However, all the methods dependent on $\beta f(\beta)$ can be directly applied to the antiferromagnetic models by transforming Jaccordingly. This means that no new HTSEs need to be calculated. We show this at the end of Sec. III E for the bcc and sc lattice. On the other hand, in the antiferromagnetic case, $\overline{\chi}(\beta)$ only presents a weak singularity if there is a finite-temperature phase transition. Because of this, it is usually better to use the HTSEs of the susceptibility associated with the magnetic order, which has to be calculated for each lattice. Except for the *fcc* and pyrochlore lattices, all the lattices mentioned above are bipartite, and thus the corresponding susceptibility is the susceptibility associated to a staggered magnetic field [8]. For the *fcc* and pyrochlore lattices, or any other nonbipartite lattice, the existence and nature of a phase transition in the antiferromagnetic case is not trivial [31–35]; and, therefore, the definition of the susceptibility related to the order parameter is more complicated. It should also be mentioned that only $\overline{\chi}(\beta)$ associated to an uniform magnetic field is accessible experimentally.

A. Dlog Padé method applied to $\overline{\chi}(\beta)$

We use first the Dlog Padé method on $\overline{\chi}(\beta)$, which is the standard method to obtain the values of T_c and the critical exponent γ from HTSEs [7,9]. The results are shown in Fig. 1 for the ferromagnetic Heisenberg model on the *fcc*, *bcc*, *sc*, pyrochlore, and *ssc* lattices. Taking into account all the poles β_i of the Padé approximants of the logarithmic derivative of $\overline{\chi}(\beta)$, we define the density of poles as a sum of Gaussian distributions,

$$N(\beta) = \sum_{i} e^{-\frac{1}{2} \left(\frac{\beta_i - \beta}{\sigma}\right)^2}$$
(14)

where $\sigma = 0.0002$ for the first three lattices and $\sigma = 0.005$ for the latter. For each lattice, we use the poles from the four highest orders in the corresponding HTSE.

The *fcc* lattice has the highest coordination number Z = 12, and the highest critical temperature (smallest β_c) of all lattices



FIG. 2. Density of residues (γ) from the Dlog Padé method on $\overline{\chi}(\beta)$ for the ferromagnetic case on the *fcc*, *bcc*, *sc*, *ssc*, and pyrochlore lattices as a function of γ . For each lattice, we use the four highest orders. The dashed line indicates the result from field theory renormalization group [14,15].

studied. As a consequence, the HTSE exploitation leads to high-quality results, even though the large Z limits the highest order n that can be reached. We can see from Fig. 1 that the values of β_c are concentrated around a well-defined value, $\beta_c = 0.4982(2)$, in agreement with the previous calculations with the same method [7,10]. For the *bcc* lattice (Z = 8) we get $\beta_c = 0.7937(2)$, and for the *sc* lattice $\beta_c = 1.1926(2)$, both of which agree with previous results [7]. For the pyrochlore lattice we obtain $\beta_c = 1.39(1)$. This last value is larger than in the sc lattice even though they both have the same coordination number. It has been argued that this is caused by the contribution of antiferromagnetic states to the partition function, which is more important in the pyrochlore than in the sc lattice [11,36]. Due to frustration, the energy difference between ferro- and antiferromagnetic states in the pyrochlore lattice is less than in the sc lattice. Finally, for the ssc lattice, the poles are too scarce and scattered to extract accurate values of β_c (lower peaks in Fig. 1 while using a higher σ). This is not surprising in the case of the *ssc*, even at orders as high as 20: because of the low coordination number Z = 3, β_c is large and the system is close to the limit Z = 2(where the system can be mapped into a 1D chain, with no singularity at finite temperatures).

From the residues, we can calculate the value of the critical exponent γ . Since all five lattices belong to the same universality class, γ is the same and thus we gather all the results in Fig. 2. In this case, we use Eq. (14) with $\beta_i \rightarrow \gamma_i$ and $\sigma = 0.004$. As can be seen in the figure, the residues from the pyrochlore and ssc lattices do not contribute significantly to the final result. In total we get $\gamma = 1.428(10)$, in agreement with previous results [7,10]. This is different from the renormalization group value, $\gamma = 1.3895(50)$ [14,15], see dashed line in Fig. 2. It was proposed that this discrepancy comes from the low order of the HTSE and that higher orders might bring the numbers closer together, as it happens for the Ising model [7,9,37]. However, the present inclusion of higher orders does not seem to point in that direction. This is indicating that a lot more orders would be needed to see an appreciable shift toward the renormalization group value.



FIG. 3. Poles and residues from the Dlog Padé method on $\overline{\chi}(\beta)$ for the ferromagnetic case on the *fcc*, *bcc*, *sc*, *ssc*, and pyrochlore lattices as a function of $(\beta - \beta_c)/\beta_c$. The inset shows the poles and residues for the pyrochlore lattice.

So far, β_c and γ have been obtained from standard methods, albeit with more orders. Alternatively, we can use our knowledge of γ to get β_c (see Fig. 3). In the Dlog Padé method, each Padé approximant provides a singularity at a given β , and its residue gives the value of the critical exponent. In practice, these couples of β and γ are not randomly scattered and, instead, fall over a monotonic increasing function. Larger values of β are accompanied by larger values of γ . In the end, for all lattices, the residues from Fig. 1 plotted versus their poles β from Fig. 2 can be plotted, and the intersection with $\gamma \simeq 1.4$ gives an approximation of β_c . Thus, with a correct choice of β_c , the residues versus $(\beta - \beta_c)/\beta_c$ should collapse on lines crossing at the universal γ for all lattices. Surprisingly, the lines of the *fcc*, *bcc*, *sc*, and pyrochlore lattices present similar slopes, whereas it is smaller for ssc. The *ssc* line gives residues at γ for $\beta_c = 4.20(5)$ (see Fig. 3). The behavior of these lines could help to determine critical values when the points do not accumulate near a single point (β_c, γ) , as in the *ssc* lattice.

Another alternative is to use the diagonal Dlog Padé method as presented in Ref. [13]. In this method, the Padé approximants of the inverse logarithmic derivative of χ are calculated, and only those with the same order in the denominator and numerator (the diagonal ones) are taken into account. Estimates of β_c are then obtained from the least positive root of the numerator (unless it is also a root of the denominator). These values converge when the order *n* increases, as illustrated for the pyrochlore and *ssc* lattices in Table II. In the case of the pyrochlore, we obtain $\beta_c = 1.39(1)$ in agreement with our previous results. However, for the *ssc* lattice we get a more precise estimate $\beta_c = 4.20(1)$.

TABLE II. Diagonal Dlog Padé results for β_c for the ferromagnetic case on the *ssc* and pyrochlore lattices.

n	8	10	12	14	16	18	20
pyrochlore ssc	1.314 5.043	1.408 4.343	1.380 4.359	1.394 4.351	1.394 4.206	4.209	4.202

B. Dlog Padé method applied to $c_v(\beta)$

So far, β_c and the critical exponent γ have been determined using the ferromagnetic susceptibility, which presents a strong singularity. The problem with relying on χ is that it depends on the order parameter, which is not generally known. Even in cases where it is known, like for antiferromagnetic models (J > 0) presenting a phase transition (bipartite lattices), new HTSEs for the antiferromagnetic staggered susceptibility χ_{AF} have to be calculated in order to see a strong singularity [8], which is computationally expensive. On the other hand, other thermodynamic functions such as the specific heat $c_v(\beta)$ and the entropy $s(\beta)$ are always indicative of a phase transition, and ferro- and antiferromagnetic models are connected by the transformation $\beta \rightarrow -\beta$. So the advantage is that no new HTSEs have to be calculated. However, there is also a disadvantage. In these functions, the ferro- and antiferromagnetic singularities coexist on the HTSE and are always present on the positive and negative β axis. Keeping this in mind, we now try to characterize a phase transition in the universality class of the Heisenberg 3D model, but without knowing the order parameter (i.e., χ): for this, we now focus on the $c_v(\beta)$ function.

Since c_v behaves as $B - A(\beta_c - \beta)^{-\alpha}$ with $-1 < \alpha < 0$ close to β_c , the Dlog Padé method cannot be used directly (the logarithmic derivative of c_v does not have a simple pole at β_c). However, the Dlog Padé method can be used on $c_v - B$. Doing so provides a good number of poles near the accepted β_c . As *B* is *a priori* unknown, we can select its value such that we get the highest quality of results and then deduce α . The issue is that there is a wide range of *B* values that give high-quality results. However, this method allows us to obtain a well-defined dependency between the height of the peak *B* and the critical exponent α . The resulting $B(\alpha)$ is displayed in Figs. 6 and 7 below, together with our interpolation method results.

C. Dlog Padé method applied to $\overline{\chi}(e)$

Finally, we can also study the singularity in $\overline{\chi}(e)$, where *e* is the energy per site. To determine the type of singularity of this function at the transition, occurring at the critical energy e_c , we start from the singularity in c_v , which can be rewritten as

$$c_v^s(\beta) = B - \tilde{A}\Delta T^{-\alpha},\tag{15}$$

where $\tilde{A} = A/T_c^{-2\alpha}$ and $\Delta T = T - T_c$. By integration, we get that close to the critical point

$$\Delta e = B\Delta T + \frac{\tilde{A}}{1-\alpha}\Delta T^{1-\alpha} + o(\Delta T^{1-\alpha}), \qquad (16)$$

where $\Delta e = e - e_c$. To get information on the singularity in $\overline{\chi}(e)$ we need the inverse $\Delta T(\Delta e)$. Since α is negative, the leading order is in ΔT . Then

$$\Delta T = \frac{\Delta e}{B} - \frac{A}{(1-\alpha)B^{2-\alpha}} \Delta e^{1-\alpha} + o(\Delta e^{1-\alpha}).$$
(17)

Keeping only the leading order and knowing that $\overline{\chi}^{s}(T) \propto \Delta T^{-\gamma}$ leads to the simple result that $\overline{\chi}^{s}(e) \propto \Delta e^{-\gamma}$, and the Dlog Padé method should give e_c as pole and γ as residue. However, taking into account that α is between -0.1 and



FIG. 4. Density of poles from the Dlog Padé method on $\overline{\chi}(e)$ for the ferromagnetic model. Results are shown as a function of $e - e_c$, where e_c is the value at which there is a peak. For each lattice, poles from HTSEs at orders from n - 3 to n are used, where n is given in Table I.

-0.2, the second leading term has similar order compared to the leading term. The quotient between both terms depends on $\Delta e^{-\alpha}$. This corresponds to a cusp-like singularity that reaches 0 only at the critical energy $\Delta e = 0$. For example, using typical values of A, B, and T_c on the fcc lattice, this quotient is about 1 when $\Delta e = 1$ and about 0.25 when $\Delta e = 0.0001$. In conclusion, only singularities at $\Delta e = e - e_c = 0$ exist, but the simple pole assumption is valid only infinitesimally close to the critical point and therefore the HTSEs are not able to represent it accurately. Thus, the Dlog Padé method can be used to obtain values for e_c from the poles, but the residues cannot capture the values of γ or α .

From the HTSEs of $\beta f(\beta)$ and of $\overline{\chi}(\beta)$ at order *n*, we obtain the series of $\overline{\chi}(e)$ at order n-1 [because the series of $e(\beta)$ are of order n-1]. Then we use the Dlog Padé method on $\overline{\chi}(e)$ and obtain the critical energies for all lattices in the ferromagnetic case (see Fig. 4). We use Eq. (14) with e (instead of β) and $\sigma = 0.01$ for the fcc, bcc, sc, and pyrochlore lattices, and $\sigma = 0.001$ for the *ssc* lattice. We find $e_c = -0.87(1)$ for the *fcc* (~58% of the ground-state energy), $e_c = -0.61(1)$ for the *bcc* (~62%), $e_c = -0.52(1)$ for the sc (~70%), $e_c = -0.57(1)$ for the pyrochlore (~76%), and $e_c = -0.302(1)$ for the ssc (~81%). Contrary to the $\overline{\chi}(\beta)$ case, the method on $\overline{\chi}(e)$ works notably better for the *ssc* lattice than for the rest. The pyrochlore lattice has a noticeably lesser amount of poles around e_c . The reason is again that this lattice is not bipartite, and the antiferromagnetic solutions on the positive e axis are frustrated. This leads to a large number of poles appearing in the positive e axis at values $e^* < |e_c|$. Finally, the residues are different for all lattices, indicating a dependency on nonuniversal quantities such as A, B, and T_c . As was expected from our previous analysis, α or γ cannot be extracted.

D. Z = 2 limit

We summarize our results obtained with the Dlog Padé method for *fcc*, *bcc*, *sc*, pyrochlore, and *ssc* lattices in Fig. 5. We plot the critical temperature T_c extracted from $\overline{\chi}(\beta)$, together with the difference between the critical energy e_c



FIG. 5. Critical temperature T_c (top panel) and the difference between the ground-state and critical energies $e_0 - e_c$ (bottom panel) as a function of the coordination number Z for the ferromagnetic model on the *fcc*, *bcc*, *sc*, *ssc*, and pyrochlore lattices obtained from the Dlog Padé method on $\overline{\chi}$. We also show the interpolation between the *sc* and *ssc* lattices by using an effective Z (see main text).

extracted from $\overline{\chi}(e)$ and the ground-state energy e_0 (known exactly for the ferromagnetic case), as a function of the coordination number Z. Both T_c and e_c show a linear behavior with respect to Z, and Z = 2 is a critical point for the finite temperature transitions in 3D ferromagnets [13], corresponding to a one-dimensional chain, characterized by $T_c = 0$ and $e_c = e_0$. To get more points, we define the *sc-ssc* model, interpolating between *sc* and *ssc* lattices (for which we have the HTSEs up to order n = 13), with two types of links on the cubic lattice

$$\mathcal{H} = J_1 \sum_{\langle ij \rangle} \mathbf{S}_i \cdot \mathbf{S}_j + J_1' \sum_{\langle ij \rangle'} \mathbf{S}_i \cdot \mathbf{S}_j$$
(18)

in such a way that when $J_1 = J'_1 = 1$ we have the *sc* lattice. When $J_1 = 1$ and $J'_1 = 0$ (or vice versa), the Hamiltonian becomes that of the *ssc* lattice. We also know that the coordination number goes from Z = 3 at $J'_1 = 0$ to Z = 6 at $J'_1 = 1$, so we can define an effective coordination number $Z_{\text{eff}}(J'_1) = 3(J_1 + J'_1)$ such that e_0 is proportional to Z_{eff} . The discrepancies between pentagons and circles of Fig. 5 at Z =3 and 6 (more visible for the energies) are due to different HTSE orders (13 for the *sc-ssc* lattice, and 17 and 20 for the *sc* and *ssc*, respectively). In addition, we also continued our calculations for the frustrated case $J'_1/J_1 < 0$ and found that T_c vanishes for $(J'_1)_c = -0.15(3)J_1$.

E. Interpolation methods for $c_v(\beta)$

The interpolation methods IM1 and IM2 presented in Sec. II B for $c_v(\beta)$ have a three dimensional parameter space: (β_c, A, α) and (β_c, B, α) for IM1 and IM2, respectively. It is one more than the similar method used for logarithmic divergencies [28], which does not have to determine a critical exponent. Exploring the whole parameter space is thus time consuming, so it is convenient to rely on other methods to narrow down some of the dimensions. In this sense, the Dlog Padé method on $\overline{\chi}$ studied in Sec. III A provides accurate values for the inverse critical temperature β_c . Thus, we leave this parameter fixed. Regarding the value of α , we know that the renormalization group value is -0.122(10) [15], while indirect estimations from HTSEs throw out values up to



FIG. 6. Top panel: Values of the singularity parameters A and B as a function of α for the ferromagnetic case on the *fcc* lattice obtained from the two interpolation methods IM1 and IM2. We show the results for the three highest orders of the HTSE of c_v . The pink dots are the results obtained with the Dlog Padé method on $c_v(\beta) - B$. Bottom panel: The quality Q [see Eq. (9)].

-0.200(15) [10]. So, for this parameter, we will search in the range [-0.05, -0.30] at 0.01 intervals. For each value of α we search for the best value of A or B (for IM1 or IM2) in a range from 0.1 to 9, at 0.002 intervals. This step must be small since the peaks in Q(A) for a given α and β_c tend to be very narrow.

Figure 6 shows the *A* and *B* values depending on the choice of α , obtained through IM1 and IM2, for the *fcc* lattice, together with the Dlog Padé method results on $c_v - B$, discussed in Sec. III B. The results for *A* and *B* show a good convergence with the HTSE order (especially at higher values of α). Furthermore, there is a good agreement between all three methods. Let us recall that IM1 and IM2 remove the singularity by subtracting and dividing, respectively, such that for IM1, *A* is a fitting parameter and *B* is a byproduct. For IM2 it is the other way around. The quality *Q* takes high values throughout all the α range: over 80% of Padé approximants coincide past β_c [up to $(1 + \delta)\beta_c$]. Even though there is a tendency to higher *Q* values for α , even choosing more restricting values for the *Q* parameters, δ , and ϵ .

For the *bcc* lattice, *Q* is between 0.7 to 0.9, whereas on the *sc* lattice, goes from 0.5 to 0.7 as α gets closer to zero. However, having half of the Padé approximants down to T_c is still a very good solution, since none of them are the same at that point when taking the raw HTSE. Also, these lattices show a convergence of the *A* and *B* values with the HTSE order *n* that is similar to the *fcc* lattice, using IM1 and IM2. *A* and *B* values for the highest HSE order are given in Fig. 7 for the *fcc*, *bcc*, and *sc* lattices. For all lattices, IM1 and IM2 give similar results, especially in the case of *B*. Differences only show up for *A* at values of α far from 0. We obtained higher values for *A* and *B* than those in the literature using HTSEs [6,7]. However, the latter were calculated by fitting the Padé approximants of the raw c_v -HTSEs with the critical behavior from Eq. (6) in a small range $\beta < 0.96 \beta_c$.



FIG. 7. Values of the nonuniversal parameters *A* and *B* as a function of α for the ferromagnetic model on the *fcc*, *bcc*, and *sc* lattices; using only the highest order in the HTSEs. Results obtained with IM1 and IM2 are shown with full and dashed lines, while symbols correspond to the Dlog Padé method on $c_v(\beta) - B$.

Another interesting feature is that the values of *B* (the peak height) are very similar for the three lattices, while the values of *A* seem to change slowly as the coordination number *Z* changes. A similar thing happens with the parameters of the singularities on Ising models on 2D, where the values are very similar but not universal in different lattices [28]. For the pyrochlore lattice, *Q* takes lower values, between 0.3 and 0.4. However, we can still extract values of *A* and *B*. They are smaller than in the *sc* lattice, even though both have the same coordination number *Z*. Taking these four lattices, *A* and *B* decrease as T_c decreases. When Z = 2 there is no finite-temperature phase transition, so it might be interesting to see how this limit is reached in terms of *A* and *B*. Finally, for the *ssc* lattice, no clear peak can be determined.

Even though it is not possible to determine the critical exponent α with these methods, we obtain well-defined functions for $A(\alpha)$ and $B(\alpha)$. Thus, we can reconstruct c_v above T_c for a supposed value of α . Using the reconstructed c_v to calculate the critical energy e_c by integration, which depends on the parameters, one could attempt to determine α by comparing with the Dlog Padé results. For the fcc lattice, the values of e_c go from -0.860(-0.857) to -0.867(-0.867) for IM1 (IM2) as α changes from -0.3 to -0.05. Again, higher values of α show a better agreement between methods. These values, summed up as $e_c = -0.862(5)$, are in agreement with the Dlog Padé estimation from the previous section $[e_c =$ -0.87(1)]. Unluckily, the Dlog Padé method does not offer sufficiently precise values of the energy and the function $e_c(\alpha)$ obtained by integration changes very little. So that in the end, it is not possible to use this extra information to determine the value of α , which remains elusive. The same happens for all the remaining lattices. For the bcc lattice we get $e_c = -0.607(3)$, in agreement with our Dlog Padé result $e_c =$ -0.61(1). For the sc lattice $e_c = -0.511(2)$, in agreement with $e_c = -0.52(1)$ from the Dlog Padé method. Finally, for the pyrochlore we get $e_c = -0.578(3)$, in agreement with $e_c = -0.57(2)$ obtained from the Dlog Padé method.



FIG. 8. Reconstructed c_v from the two interpolation methods for the ferromagnetic case on the *fcc*, *bcc*, *sc*, and pyrochlore lattices. We show the results for two different values of α . The inset shows a zoom for the *fcc* lattice close to the critical temperature.

The critical entropy can also be obtained by integration. Let us start with the *sc* lattice, for which we obtain $s_c = 0.402(1)$, where $s_c = 0.401$ for $\alpha = -0.1$ and $s_c = 0.403$ for $\alpha = -0.2$ with both interpolation methods. In this case, it is possible to benchmark with QMC results, which give $s_c = 0.401(5)$ [38]. All of our s_c values are in agreement with QMC while showing a slightly better precision. Also, it is not possible to decide on α from this calculation. For the *bcc* lattice, all of our results are within $s_c = 0.435(1)$. For the *fcc* lattice we get $s_c = 0.443(1)$. Finally, for the pyrochlore, we obtain $s_c = 0.353(2)$. All in all, we can see that the critical entropy decreases with the coordination number, as does the critical temperature. However, contrary to the results shown in Fig. 5 for the critical temperature and $e_0 - e_c$, the critical entropy s_c does not show a linear behavior towards 0 at Z = 2.

We show in Fig. 8 the reconstructed c_v for the four lattices (fcc, bcc, sc, and pyrochlore) using the best values of A and B at the accepted β_c for two limiting values of α , -0.2 and -0.1. Both methods give the same curves, and the differences between the two values of α can only be seen very close to the corresponding critical points (see inset for fcc) through a very different value of the peak height B, as can be seen in the previous figures of $B(\alpha)$. However, this issue only exists at exactly the critical temperature, so it does not affect the comparison with experimental results since the sharp theoretical peaks with divergent slopes at the critical point cannot be captured by experiments in real compounds [18,39–42]. To sum up, we have a good precision for every temperature above the critical temperature T_c obtained from finite high-temperature series expansions. Thus, this method extrapolates the specific heat from HTSE down to almost the critical temperature for the phase transitions of several ferromagnetic Heisenberg models.

As we mentioned earlier, the advantage of using c_v instead of χ resides in the possibility of studying both ferro- and antiferromagnetic models with the same HTSE. In Fig. 9 we show results obtained for the antiferromagnetic *sc* and *bcc* lattices. For these calculations, since one wants to avoid using



FIG. 9. Interpolation method results for $c_v(\beta)$ for the antiferromagnetic model on the *sc* (left) and *bcc* (right) lattices. The quality of results *Q* is shown in color scale in a region of the parameter space defined by T_c and *A* or *B*. The critical exponent α is set to -0.12. The quality *Q* as a function of T_c is shown in the top panels, in blue and orange for IM1 and IM2, respectively. Dashed black lines indicate QMC results [43,44].

data for $\overline{\chi}(\beta)$, we calculated T_c by fixing $\alpha = -0.12$, close to the field theory result. However, the critical temperature is not very sensitive to this constraint, at least within the range of values reported in the literature. In the top panels of Fig. 9 we show the values of Q as a function of T_c obtained from the color plots. For both lattices, there is a well-defined region of parameters with high Q, meaning that almost all Padé approximants coincide down to the critical temperature. Also, these regions of high Q agree well with QMC results, indicated by the dashed black lines [43,44]. For the sc lattice we get $T_c = 0.93(2)$, in agreement with the QMC result $T_c = 0.946(1)$ [43]. For the *bcc* lattice we get 1.38(4), in agreement with the QMC result $T_c = 1.377(2)$ [44]. We can also take the best values from each interpolation method to reconstruct c_v for the antiferromagnetic cases and calculate the critical energies and entropies. For the sc lattice, we get $e_c = -0.66(1)$ and $s_c = 0.331(5)$. The latter is slightly lower than the QMC result, $s_c = 0.341(5)$ [38], but still fairly close. For the *bcc* lattice we get $e_c = -0.73(1)$ and $s_c = 0.393(5)$.

F. Interpolation method for $\chi(\beta)$

Finally, we apply the interpolation method IM2 to the ferromagnetic $\overline{\chi}(\beta)$ (as explained in Sec. II C for divergent singularities). The parameter space is two-dimensional $\{\beta_c, \gamma\}$, and the region of high-quality values is narrow so both parameters have to be calculated using a fine mesh.

Figure 10 shows the quality Q as a function of β_c and γ for the *fcc*, *bcc*, *sc*, and pyrochlore lattices. We also show in white circles the results from the poles and residues obtained from the Dlog Padé method. For all lattices, the poles and residues are concentrated around the large Q region from IM2, and reciprocally, the higher Q values are obtained close to the line of poles and residues from the Dlog Padé method.



FIG. 10. Interpolation method IM2 results for $\overline{\chi}(\beta)$ for the ferromagnetic case on the *fcc*, *bcc*, *sc*, and pyrochlore lattices. The quality of results *Q* is shown in color scale in a region of the parameter space $\{\beta_c, \gamma\}$ close to the best values. White circles indicate the poles and residues from the Dlog Padé method.

This illustrates a close connection between both methods. Specifically, the *fcc* lattice presents Q = 1.00 around $\beta_c = 0.4981(2)$ and $\gamma = 1.422(2)$, the *bcc* lattice presents Q = 0.93 for $\beta_c = 0.7938(2)$ and $\gamma = 1.420(3)$, the *sc* lattice presents Q = 0.64 for $\beta_c = 1.1935(10)$ and $\gamma = 1.44(1)$. All of these are in agreement with the Dlog Padé results. For the pyrochlore and *ssc* (not shown) lattices, the results are not so clear. The pyrochlore lattice has a large cloud of values Q = 0.41(1) along a well-defined line around $\beta_c = 1.382(5)$ and $\gamma = 1.36(3)$. However, the diagonal Dlog Padé results lie closer to the endpoint of this cloud. For the *ssc* lattice, there are just a few points around Q = 0.23(2) with $\beta_c = 4.20(2)$ and $\gamma = 1.35(2)$, but the quality is too low to consider them reliable.

IV. CONCLUSIONS AND PERSPECTIVES

We have studied the finite-temperature phase transition that occurs in ferromagnetic quantum Heisenberg models on 3D lattices by using several methods derived from the HTSEs. We used the standard Dlog Padé method and estimated β_c and γ of the *fcc*, *bcc*, *sc*, pyrochlore, and *ssc* lattices. For some of them, results are given for larger orders of HTSEs than in the previous papers. However, the discrepancy in γ between field theory's renormalization group for the classical case and the HTSEs results for the quantum case is still present. Also, no convergence towards the classical values is observed, and this remains an open question. We have also explored possible extensions of these methods. While standard calculations involve $\chi(\beta)$, we have obtained the critical energy e_c using $\chi(e)$. We also used the Dlog Padé method on $c_v(\beta) - B$ [with $B = c_v(\beta_c)$] to obtain $B(\alpha)$.

Then we presented interpolation methods to obtain $c_v(T)$ and $\chi(T)$ for $T > T_c$. These methods are efficient for the *fcc*,

TABLE III. Summary of results for β_c , e_c , and s_c obtained in this article for the ferromagnetic Heisenberg model on the *fcc*, *bcc*, *sc*, pyrochlore (pyro), and *ssc* lattices. DLP stands for Dlog Padé method and IM for the interpolation method.

	eta_c		e	S _c	
Lattice	DLP	IM	DLP	IM	IM
fcc	0.4982(2)	0.4981(2)	-0.87(1)	-0.862(5)	0.443(1)
bcc	0.7937(2)	0.7938(2)	-0.61(1)	-0.607(3)	0.435(1)
sc pyro ssc	1.1926(2) 1.39(1) 4.20(1)	1.1935(10) 1.382(5) 4.20(2)	-0.52(1) -0.57(2) -0.302(1)	-0.511(2) -0.578(3)	0.402(1) 0.353(2)

bcc, and *sc* lattices, but less for the pyrochlore and *ssc* lattices. For $c_v(T)$, we are not able to get a precise value of the critical exponent α , but the methods provide accurate relationships between the three important parameters at the singularity, A, B, and α . Thus, if any of them is known, the other two can be deduced. We have also shown that the interpolated $c_v(T)$ has a very small dependence on α as soon as T is slightly above T_c , the main difference being in the value of the peak at T_c . This allowed us to obtain accurate results of the critical energy e_c and critical entropy s_c for most lattices studied. The main advantage of the method is that it can be directly applied to antiferromagnetic cases without the need of calculating a new HTSE. To show this, we applied both interpolation methods to the bcc and sc lattices, and obtained results in agreement with QMC for T_c and s_c . Finally, we applied the interpolation method to obtain $\chi(T)$ above T_c, obtaining reliable values of T_c and γ .

In conclusion, we have probed several different methods based on HTSEs to study finite-temperature phase transitions. These methods allowed us to obtain accurately several quantities related to the critical points, such as critical exponents, critical temperatures, and parameters related to the singularities. As a summary, we present the main numerical results in Tables III and IV, where DLP stands for the Dlog Padé results and IM stands for the interpolation method. The results shown for *A* are obtained with IM, while the ones for *B* cover both IM and Dlog Padé results. It is important to note that even if we have approached only the ferro- and antiferromagnetic Heisenberg models on the most common lattices without frustration, these methods are suitable for studying any kind of system with the same type of phase transitions.

TABLE IV. Summary of results for γ , A, and B obtained in this article for the ferromagnetic Heisenberg model on the *fcc*, *bcc*, *sc*, pyrochlore (pyro), and *ssc* lattices. DLP stands for Dlog Padé method and IM for the interpolation method.

	γ		$\alpha = -0.1$		$\alpha = -0.2$	
Lattice	DLP	IM	Α	В	Α	В
fcc	1.426(2)	1.422(2)	3.25(5)	2.90(5)	2.41(5)	1.74(3)
bcc	1.419(2)	1.420(3)	3.10(5)	2.85(5)	2.17(6)	1.71(2)
sc	1.433(3)	1.44(1)	2.71(3)	2.62(5)	1.80(3)	1.56(3)
pyro ssc		1.36(3) 1.35(2)	2.15(15)	2.1(1)	1.33(3)	1.22(3)

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APPENDIX: HTSEs FOR THE S = 1/2 MODELS

In Tables V–VIII we present the complete list of coefficients for the HTSEs of βf and $\overline{\chi}$, where the new ones are in bold numbers. These are written in terms of Eq. (3) where $n_u = 1$ for the *fcc*, *bcc*, and *sc* lattices; $n_u = 4$ for the *ssc* and pyrochlore lattices.

TABLE V. Coefficients a_n corresponding to βf as defined in Eq. (3) for the *fcc*, *bcc*, and *sc* lattices.

a_n	fcc	bcc	SC
1	0	0	0
2	18	12	9
3	-108	24	18
4	180	168	-162
5	5040	-1440	-2520
6	162000	24480	33192
7	-14565600	297024	1019088
8	563253408	28017216	-7804944
9	-17544639744	533681664	-723961728
10	750412309248	41156316672	2596523904
11	-56646776913408	503287538688	856142090496
12	4973976625190400	53001415916544	6383648984832
13	-421817449494804480	1839416689004544	-1356696930401280
14		246102905022713856	-27667884260938752
15		9001661201883684864	2908030732698175488
16			122264703581556307968
17			-7238339805811283361792

TABLE VI. Coefficients b_n corresponding to $\overline{\chi}$ as defined in Eq. (3) for the *fcc*, *bcc*, and *sc* lattices.

b_n	fcc	bcc	SC
1	-6	-4	-3
2	120	48	24
3	-3312	-832	-264
4	117360	18400	3960
5	-5104416	-504384	-74928
6	263405088	16313280	1584624
7	-15717292800	-610699520	-38523264
8	1063892512512	25867292160	1115604864
9	-80532234584064	-1229543182336	-35969253888
10	6741740335372800	64541249655808	1223162767104
11	-618536855295817728	-3716345369001984	-46443134693376
12	61718837768472705024	232442811396567040	1997899947119616
13	-6654017125619285385216	-15728092831910068224	-90256082576916480
14		1142407619109235630080	4221503453720782848
15		-88805626440393148956672	-220236945885669801984
16			12562562473105938481152
17			-722105535259151290073088

a_n	pyrochlore	SSC
1	0	0
2	36	18
3	-72	36
4	-1656	-324
5	18720	-3600
6	340704	20592
7	-11342016	788256
8	-99460800	-267552
9	11144157696	-292582656
10	-43247015424	-2338428672
11	-15542133488640	158857763328
12	359762974166016	3398565523968
13	26719647518453760	-111431579830272
14	-1532961802218000384	-5116416515020800
15	-44591351194841260032	83476825611595776
16	6653104879154138357760	9038092645962510336
17	-8491904138175731859456	-20724045060052942848
18		-18839898190998133604352
19		-253691725481243238334464
20		45155117370822689756676096

TABLE VII. Coefficients a_n corresponding to βf as defined in Eq. (3) for the pyrochlore and *ssc* lattices.

TABLE VIII. Coefficients b_n corresponding to $\overline{\chi}$ as defined in Eq. (3) for the pyrochlore and *ssc* lattices.

$\overline{b_n}$	pyrochlore	SSC
1	-12	-6
2	96	12
3	-816	12
4	8160	240
5	-148992	-5136
6	3879744	-40224
7	-81019776	778464
8	990764544	22859520
9	-15165570048	-183876864
10	1661765784576	-15637820928
11	-97979429505024	-30648860160
12	1761563239919616	14373541840896
13	85410304429842432	215523347675136
14	-1996581576629084160	-16345519886733312
15	-507546664875986436096	-572943639086174208
16	35052604281755859025920	20821759189681766400
17		1524898473896350777344
18		-22745675831893506785280
19		-4446640583932914089459712
20		-17616386456676250248806400

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