5 Dynamical Vertex Approximation

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Contents

1 Introduction 2

2 Feynman diagrammatics 3
  2.1 Parquet equations 3
  2.2 Dynamical vertex approximation (DΓA) 10

3 Two highlights 15
  3.1 Critical exponents of the Hubbard model 15
  3.2 Fate of the false Mott-Hubbard transition in two dimensions 17

4 Conclusion and outlook 19
1 Introduction

The theoretical description and understanding of strongly correlated systems is particularly challenging since perturbation theory in terms of the Coulomb interaction is no longer possible and standard mean field theory does not work. Also bandstructure calculations in the local density approximation (LDA) which had been so successful for the calculations of many materials do not work properly as electronic correlations are only rudimentarily taken into account. A big step forward in this respect is dynamical mean field theory (DMFT) which is a mean field theory in the spatial coordinates, but fully accounts for the local correlations in time (quantum fluctuations). In comparison, standard Hartree-Fock is mean-field in space and time. DMFT is non-perturbative since all Feynman diagrams are taken into account, albeit only their local contribution for the self energy. If one is dealing with well localized d- or f-orbitals, these local DMFT correlations often provide the major part of the electronic correlations, which not only give rise to mass renormalizations, metal-insulator transitions and magnetic ordering, but also to unexpected new physics such as kinks or the filling of the Mott-Hubbard gap with increasing temperature. More aspects of DMFT are discussed in other contributions to this Jülich Autumn School on DMFT at 25.

The question we would like to address here is: Can we do (or do we need to do) better than DMFT? Indeed, going beyond DMFT is necessary since many of the most fascinating and least understood physical phenomena such as quantum criticality and superconductivity originate from non-local correlations – which are by construction beyond DMFT. And we can: A first step to include non-local correlations beyond DMFT have been cluster approaches such as the dynamical cluster approximation (DCA) and cluster DMFT (CDMFT). Here, instead of considering a single site embedded in a mean-field medium. Numerical limitations however restrict the DCA and CDMFT calculations to about 100 sites. This allows for studying short-range correlations, particularly for two-dimensional lattices, but severely restricts the approach for three dimensions, for multi-orbitals, and regarding long-range correlations.

Because of that, in recent years diagrammatic extensions of DMFT were at the focus of the methodological development. An early such extension was the 1/d (d: dimension) approach: also the combination of the non-local spin fermion self energy with the local quantum fluctuations of DMFT has been proposed. Most actively pursued are nowadays diagrammatic approaches based on the local two-particle vertex. The first such approach has been the dynamical vertex approximation (DΓA), followed by the dual fermion approach, the one-particle irreducible approach (1PI), and DMFT to functional renormalization group (DMF2RG).

The very idea of these approaches is to extend the DMFT concept of taking all (local) Feynman diagrams for the one-particle irreducible vertex (i.e., the self energy), to the next, i.e., two-particle level. In these approaches, one calculates the local two-particle vertex; and from this non-local correlations beyond DMFT are obtained diagrammatically. Indeed, we understand most (if not all) physical phenomena either on the one-particle level (e.g. the quasiparticle renor-
Table 1: Summary of the different diagrammatic extensions of DMFT based on the two-particle vertex. All methods are based on the local part of the two-particle vertex named in the table; for a definition of the different vertex functions, see Section 2.1.

| Method         | Local two-particle vertex | Feynman diagrams |
|----------------|----------------------------|------------------|
| DF [18]        | one-particle reducible vertex, here* \( F_{\text{loc}} \) | 2nd order, ladder parquet |
| 1PI [19]       | one-particle irreducible vertex \( F_{\text{loc}} \) | ladder          |
| DMF\(^2\)RG [20] | one-particle irreducible vertex \( F_{\text{loc}} \) | RG flow         |
| ladder D\(\Gamma\)A [17] | two-particle irreducible vertex in channel \( r \Gamma_{\text{loc}} \) | ladder          |
| full D\(\Gamma\)A [17] | two-particle fully irreducible vertex \( \Lambda_{\text{loc}} \) | parquet         |

* Note that at the two-particle level every two-particle vertex is one-particle irreducible; three or higher order vertices can be however one-particle reducible which has consequences for the diagrammatics if truncated at the two-particle level, see [19].

malization and the Mott-Hubbard transition) or on the two-particle level [e.g. (para)magnons and (quantum) critical fluctuations]. Non-local correlations and associated physics on this two-particle are included in these diagrammatic extensions of DMFT, which however still include the local DMFT one-particle physics such as the formation of Hubbard bands and the metal-insulator transition or, more precisely, a renormalized version thereof. The concept of all these approaches is similar, but they differ in which two-particle vertex is taken and which diagrams are constructed, see Table 1. Depending on the approach, Feynman diagrams are constructed from full Green function lines \( G(\nu, k) \) or from the difference between \( G(\nu, k) \) and the local Green function \( G_{\text{loc}}(\nu) \) \([\nu]:\) (Matsubara) frequency; \( k\): wave vector]. The DF, 1PI and DMF\(^2\)RG approach are also based on a generating functional integral.

In these lecture notes, we will concentrate on D\(\Gamma\)A in the following. Section 2.1 recapitulates the concept of reducible and irreducible diagrams, the parquet and Bethe Salpter equation. On this basis, we introduce in Section 2.2 the D\(\Gamma\)A approach. In Section 3 we have chosen two exemplary highlights that demonstrate what can be calculated by D\(\Gamma\)A and related approaches. These are: the calculation of the critical exponents for the three dimensional Hubbard model (Section 3.1) and the effect of long-range correlations on the Mott-Hubbard transition for the two dimensional Hubbard model (Section 3.2), at zero temperature antiferromagnetic fluctuations always open a gap at any interaction \( U > 0 \).

2 Feynman diagrammatics

2.1 Parquet equations

The very idea of D\(\Gamma\)A is a resummation of Feynman diagrams, not in orders of the interaction \( U \) as in perturbation theory but in terms of the locality of diagrams. In this sense, DMFT is the first, one-particle level since it approximates the one-particle fully irreducible vertex, i.e., the self energy \( \Sigma \), to be local, see Table 2.
Table 2: \(D \Gamma A\) generalizes the DMFT concept of the local self energy (i.e., the local fully irreducible one-particle vertex) to the fully irreducible \(n\)-particle vertex. It is hence a resummation of Feynman diagrams in terms of their locality. On the right hand side the different levels of approximation are indicated.

| \(n\) | DMFT: local self energy | \(D \Gamma A\): local fully irreducible two-particle vertex | ⇒ non-local self energy/correlations | \(n \rightarrow \infty\) exact solution |
|---|---|---|---|---|
| 1 | | | | |
| 2 | | | | |
| ... | | | | |
| \(n \rightarrow \infty\) | | | | |

Fig. 1: Basic objects of Feynman diagrams: from the non-interacting Green function \(G_0\) (left) and the bare interaction \(U\) (middle) we construct all topologically distinct diagrams for calculating the interacting Green function \(G\) (right).

For a better understanding of reducibility and irreducibility as well as of \(D \Gamma A\) later on, let us recall that in quantum field theory we calculate the interacting Green function \(G\) by drawing all topologically distinct Feynman diagrams that consist of \(n\) interactions \(U\) and that are connected by non-interacting Green function lines \(G_0\), keeping one incoming and one outgoing \(G_0\) line, see Fig. 1. Each \(G_0\) line contributes a factor \(G_0(\nu, k) = 1/(\nu + \mu - \epsilon_k)\) [where \(\nu\) denotes the (Matsubara) frequency, \(\mu\) the chemical potential and \(\epsilon_k\) the energy-momentum dispersion relation of the non-interacting problem] and each interaction (wiggled line) contributes a factor \(U\). Here and in the following, we assume a one band model for the sake of simplicity. For an introduction to Feynman diagrams, more details and how to evaluate Feynman diagrams including the proper prefactor, we refer the reader to textbooks of quantum field theory such as [21]; a more detailed presentation including \(D \Gamma A\) can also be found in [22].

**Dyson equation and self energy**

Instead of focusing on the Green function, we can consider a more compact object, the self energy \(\Sigma\), which is related to \(G\) through the Dyson equation, see Fig. 2. The Dyson equation

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1 For a \(k\)-dependent, i.e. non-local interaction the factor \(U(k, k', k'', k''') = k + k' - k''\) would be \(k\)-dependent.
Fig. 2: Dyson equation connecting the Green function and self energy. The pair of scissors indicates that these diagrams are one-particle reducible (i.e., cutting one $G_0$ line separates the Feynman diagram into two parts)

can be resolved for the interacting Green function:

$$G(\nu, \mathbf{k}) = \left[ 1/G_0(\nu, \mathbf{k}) - \Sigma(\nu, \mathbf{k}) \right]^{-1}. \quad (1)$$

Since the geometric series of the the Dyson equation generates a series of Feynman diagrams, we can only include a reduced subset of Feynman diagrams when evaluating the self energy. One obvious point is that the two outer “legs” (incoming and outgoing $G_0$ lines) are explicitly added when going from the self energy to the Green function, see Fig. 2. Hence we have to “amputate” (omit) these outer “legs” for self energy diagrams. More importantly, the self energy can only include one-particle irreducible diagrams. Here, one-particle (ir)reducible means that by cutting one Green function line one can(not) separate the diagram into two parts. This is since, otherwise, we would generate Feynman diagrams twice: Any one-particle reducible diagram can be constructed from two (or more) irreducible building blocks connected by one (or more) single $G_0$ lines. This is exactly what the Dyson equation does, see Fig. 2. For example, in the last line, we have three irreducible self energy blocks connected by two single $G_0$ lines. This shows that, by construction, the self energy has to include all one-particle irreducible Feynman diagrams and no one-particle reducible diagram. Since the self energy has one (amputated) incoming leg, it is a one particle vertex. It is also one-particle irreducible as explained above. Hence, the self energy is the one-particle irreducible one-particle vertex. In Fig. 3 we show some diagrams that are part of the self energy, i.e., cutting one-line does not separate the diagram into two parts, and some diagrams that are not.

Two-particle irreducibility

Let us now turn to the next, the two-particle level. As illustrated in Fig. 4 the Feynman diagrams for the susceptibility $\chi$ (or similarly for the two particle Green function) consist of (i) an unconnected part (two $G$ lines, as in the non-interacting case) and (ii) all connected Feynman diagrams (coined vertex corrections). Mathematically this yields ($\beta$: inverse temperature; $\sigma$:}
Fig. 3: Left: some examples of (one-particle irreducible) self energy diagrams. Right: Diagrams that do no contribute to the self energy since they are one-particle reducible (cutting the line indicated by the pair of scissors separates the diagram into two pieces).

\[
\chi_{\sigma\sigma'}(\nu\nu'; k, k', q) = -\beta G(\nu, k)G(\nu + \omega, k + q)\delta_{\nu\nu'}\delta(k - k')\delta_{\sigma\sigma'} \\
+ G(\nu, k)G(\nu + \omega, k + q)F_{\sigma\sigma'}(\nu\nu'; k, k', q)G(\nu', k')G(\nu' + \omega, k' + q).
\]

Here \( F \) denotes the full, reducible vertex. In the following let us introduce a short-hand notation for the sake of simplicity, where 1 represents a momentum-frequency-spin coordinate \( (k, \nu, \sigma) \), 2 \( (k + q, \nu + \omega, \sigma) \), 3 \( (k' + q, \nu' + \omega, \sigma') \), 4 \( (k', \nu', \sigma') \). In this notation we have

\[
\chi(1234) = -G(14)G(23) - G(1'1')G(2'2')F(1'2'3'4')G(3'3')G(4'4'),
\]

as visualized in Fig. 4. Since the Green function is diagonal in spin, momentum and frequency, i.e., \( G(11') = G(11')\delta_{11'} \), some indices are the same which yields Eq. (3).

Let us now again introduce the concept of irreducibility, this time for the two-particle vertex. In this case, we consider two-particle irreducibility. In analogy to the self energy, we define the fully irreducible vertex \( \Lambda \), defined as the set of all Feynman diagrams that do not split into two parts by cutting two \( G \) lines. Let us remark that here and in the following, we construct the Feynman diagrams in terms of \( G \) instead of \( G_0 \). This means that we have to exclude all diagrams which contain a structure as generated by the Dyson equation in Fig. 2 since otherwise these diagrams would be counted twice. This reduced set of diagrams with \( G \) instead of \( G_0 \) is called skeleton diagrams, see [21].

The reducible diagrams of \( F \) can be further classified according to how the Feynman diagram separates when cutting two internal Green functions. Since \( F \) has four (“amputated”) legs to the outside, there are actually three possibilities to split \( F \) into two parts by cutting two \( G \) lines.

Note, that one-particle irreducibility is somehow trivial since one can show that there are no one-particle irreducible diagrams for the two-particle vertex (in terms of the interacting \( G/skeleton \) diagrams).
In analogy to the Dyson equation, Fig. 2, the reducible vertices are defined as reducible diagrams in Fig. 4, which consists of two unconnected Green function $G$ lines (aka “bubble”) and vertex $F$ corrections.

\[
\chi_{1234} = -\chi_{1234} - 2_{1234} + F_{34}.
\]

**Fig. 4:** The susceptibility $\chi$ consists of two unconnected Green function $G$ lines (aka “bubble”) and vertex $F$ corrections.

That is, an external leg, say 1, stays connected with one out of the three remaining external legs but is disconnected from the other two, see Fig. 5 for an illustration. One can show by hands of the diagrammatic topology, that each diagram is either fully irreducible or reducible in exactly one channel, so that

\[
F(1234) = \Lambda(1234) + \Phi_{ph}(1234) + \Phi_{pp}(1234) + \Phi_{pp}(1234).
\]

**Fig. 5:** The full (reducible) vertex $F$, consist of the fully irreducible vertex $\Lambda$ and two-particle reducible diagrams. These can be classified into three channels depending on which parts are disconnected by cutting two Green function lines. These are commonly denoted as the particle-hole reducible channel $\Phi_{ph}$ separating 12 from 34, the transversal particle-hole reducible channel $\Phi_{ph}$ separating 14 from 23, and the particle-particle reducible channel $\Phi_{pp}$ separating 13 from 24. Each two-particle reducible diagram is reducible in one (and only one) of these three channels. The hatched blocks themselves can be irreducible, reducible in the same channel*, or reducible in the other two channels (in this last case the full diagram remains however reducible only in the scissors-induced channel).

*:Note, this is only possible for one hatched side, since otherwise the same diagram might be counted twice.

\[
F(1234) = \Lambda(1234) + \Phi_{ph}(1234) + \Phi_{pp}(1234) + \Phi_{pp}(1234).
\]

**Bethe-Salpeter equation**

We have defined the reducible diagrams $\Phi_r$ in channel $r \in \{ph, ph, pp\}$ as a subset of Feynman diagrams for $F$. The rest, i.e., $F - \Phi_r$, is called the vertex $\Gamma_r$ irreducible in $r$ so that

\[
F(1234) = \Gamma_r(1234) + \Phi_r(1234) .
\]

In analogy to the Dyson equation, Fig. 2 the reducible vertices $\Phi_r$ in turn can be constructed from $\Gamma_r$. One $\Gamma_r$ can be connected by two $G$s with another $\Gamma_r$ (which makes this diagram...
Parquet equations

Since an irreducible $\Gamma_r$ diagram in a channel $r$ is either fully irreducible ($\Lambda$) or reducible in one of the two other channels $r' \neq r$ ($\Phi_{r'}$), we can express $\Gamma_r$ as [this also follows directly from Eqs. (4) and (5)]:

$$\Gamma_r(1234) = \Lambda(1234) + \sum_{r' \neq r} \Phi_{r'}(1234).$$

(9)

We can use this Eq. (9) to substitute the last $\Gamma_r$’s in Eq. (6) (or the $\Gamma_r$ box in Fig. 6) by $\Lambda$ and $\Phi_{r'}$’s. Bringing the first $\Gamma_r$ on the left hand side, then yields:

$$\Phi_{ph}(1234) = F(1234) - \Gamma_{ph}(1234)$$

$$= F(122'1')G(3'2')G(1'4')\Lambda(4'3'34) + \sum_{r' \neq \text{ph}} F(122'1')G(3'2')G(1'4')\Phi_{r'}(4'3'34)$$

(10)
Fig. 7: Parquet equations. From the Bethe-Salpeter equations in the three channels we obtain
three corresponding equations connecting the reducible vertex $F$, the fully irreducible vertex $\Lambda$
and the reducible vertices $\Phi_r$ in the three channels $r$ (first three lines). Together with the clas-
sification of $F$ into $\Lambda$ and the three $\Phi_r$ (last line) we have the four so-called parquet equations.

and two corresponding equations for the other two channels. The corresponding Feynman di-
agrams are shown in Fig. 7. If the fully irreducible vertex $\Lambda$ is known, these three equations
together with Eq. (4) allow us to calculate the four unknown vertex functions $\Phi_r$ and $F$, see
Fig. 7. This set of equations is called the parquet equations. The solution can be done nu-
merically by iterating these four parquet equations. Reflecting how we arrived at the parquet
equations, the reader will realize that the parquet equations are nothing but a classification of
Feynman diagrams into fully irreducible diagrams and diagrams reducible in the three channels
$r$.

The thoughtful reader will have also noticed that the interacting Green function enters in Eq. (10).
This $G$ can be calculated by one additional layer of self consistency. If the reducible vertex $F$
is known, the self energy follows from the Heisenberg Eq. of motion (also called Schwinger-
Dyson Eq. in this context). This is illustrated in Fig. 8 and mathematically reads:

$$
\Sigma(14) = -U(12'3'1')G(1'4')G(23')G(2'3')F(4'234) + U(1234)G(23) - U(1432)G(23)
$$

That is the numerical solution of the four parquet equations has to be be supplemented by the
Schwinger-Dyson Eq. (11) and the Dyson Eq. (1), so that also $G$ and $\Sigma$ are calculated self-
consistently.

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3 Sometimes, also only Eq. (5) is called parquet equation and the equations of type Eq. (10) remain under the
name Bethe-Salpeter equations.
Fig. 8: Equation of motion (Schwinger-Dyson equation) for calculating the self energy $\Sigma$ from the bare interaction $U$, the reducible vertex $F$ and the Green function lines $G$. The second and third diagram on the left hand side are the Hartree and Fock diagrams, respectively, which are not included in the first term.

Let us also note that these general equations while having a simple structure in the 1234 notion can be further reduced for practical calculations: The Green functions are diagonal $G(3'2') = G(3'3')\delta_{2'3'}$, there is a severe restriction in spin, there is SU(2) symmetry and one can decouple the equations into charge(spin) channels $\Gamma_{ph\uparrow\uparrow} + (-)\Gamma_{ph\uparrow\downarrow}$. A detailed discussion hereof is beyond the scope of these lecture notes. For more details on the parquet equations see [23], for a derivation of the equation of motion also see [24].

2.2 Dynamical vertex approximation ($D\Gamma A$)

Hitherto, everything has been exact. If we know the exact fully irreducible vertex $\Lambda$, we can calculate through the parquet equations Fig. 7 [Eqs. (5) and (10)] the full vertex $F$; from this through the Schwinger-Dyson equation of motion (11) the self energy $\Sigma$ and through the Dyson Eq. (1) the Green function $G$. With a new $G$ we can (at fixed) $\Lambda$ recalculate $F$ etc. until convergence. Likewise if we know the exact irreducible vertex $\Gamma_r$ in one channel $r$, we can calculate $F$ through the corresponding Bethe-Salpeter Eq. (6-8) and from this (self consistently) $\Sigma$ and $G$.

But $\Lambda$ (or $\Gamma_r$) still consists of an infinite set of Feynman diagrams which we usually do not know. Since the parquet (or Bethe-Salpeter) equations generate many additional diagrams, there are however much less (albeit still infinitely many) diagrams for $\Lambda$ (or $\Gamma_r$) than for $F$. In the case of $\Lambda$, the bare interaction $U$ is included but the next term is already a diagram of fourth (!) order in $U$ (the so-called envelope diagram), see Fig. 9. There are no two-particle fully irreducible diagrams of second or third order in $U$.

One approach is hence to approximate $\Lambda$ by the bare interaction $U$ only, i.e., the first two terms in Fig. 9. This is the so-called parquet approximation [23]. For strongly correlated electrons this will be not enough. A deficiency is, for example, that the parquet approximation does not yield Hubbard bands.

In $D\Gamma A$, we hence take instead all Feynman diagrams for $\Lambda$ but restrict ourselves to their local
contribution, \( A_{\text{loc}} \). This approach is non-perturbative in the local interaction \( U \). It is putting the DMFT concept of locality to the next, i.e., to the two-particle level. We can extend this concept to the \( n \)-particle fully irreducible vertex, so that by increasing \( n \) systematically more and more Feynman diagrammatic contributions are generated; and for \( n \to \infty \) the exact solution is recovered, see Table 2 above.

In practice, one has to truncate this scheme at some \( n \), hitherto at the two-particle vertex level \((n = 2)\). The local fully irreducible two-particle vertex \( \Lambda \) can be calculated by solving an Anderson impurity model which has the same local \( U \) and the same Green function \( G \). This is because such an Anderson impurity model yields exactly the same (local) Feynman diagrams \( \Lambda_{\text{loc}} \). It is important to note that the locality for \( \Lambda \) is much better fulfilled than that for \( \Sigma \). Even in two dimensions, \( \Lambda \) is essentially \( k \)-independent, i.e., local. This has been demonstrated by numerical calculations for the two-band Hubbard model, see [25]. In contrast, for the same set of parameters \( \Sigma \) is strongly \( k \)-dependent, i.e., non-local. Also \( \Gamma_r \) and \( F \) are much less local than \( \Lambda \), see [25]. There might be parameter regions in two dimensions or one-dimensional models where also \( \Lambda \) exhibits a sizable non-local contributions. One should keep in mind that \( D \Gamma A \) at the \( n = 2 \) level is still an approximation. This approximation includes however not only DMFT but on top of that also non-local correlations on all length scales so that important physical phenomena can be described; and even in two dimensions substituting \( \Lambda \) by its local contribution \( \Lambda_{\text{loc}} \) is a good approximation, better than replacing \( \Sigma \) by its local contribution \( \Sigma_{\text{loc}} \) as in DMFT.

When solving the Anderson impurity model numerically, one does not obtain \( A_{\text{loc}} \) directly but first the local susceptibilities \( \chi_{\text{loc}} \) (or the two-particle Green function). Going from here to \( A_{\text{loc}} \) is possible as follows: From \( \chi_{\text{loc}} \) and \( G_{\text{loc}} \), we obtain the local reducible vertex \( F_{\text{loc}} \) [via the local version of Eq. (3)], from this in turn we get \( \Gamma_{r\text{loc}} \) [via inverting the local version of the Bethe-Salpeter Eqs. (6-8)], \( \Phi_{r\text{loc}} \) [via the local version of Eq. (5)], and finally \( \Lambda_{\text{loc}} \) [via the local version of Eq. (4)].

Fig. 9: \textit{Lowest order Feynman diagrams for the fully irreducible vertex} \( \Lambda \).
Fig. 10: Top: Local full vertex $F$ in the magnetic (m) and charge or density (d) channel, i.e., $F_{\uparrow\uparrow} \pm F_{\uparrow\downarrow}$, as a function of the two incoming Fermionic frequencies $\nu = (2n + 1)\pi T$ and $\nu' = (2n' + 1)\pi T$ at transferred Bosonic frequency $\omega = 0$ for the three dimensional Hubbard model at $U = 0.5, T = 1/26$ (in units of nearest neighbor hopping $2\sqrt{6}t \equiv 1$). Middle: Corresponding particle-hole irreducible vertex $\Gamma_{ph}$ (two left panels) and particle-particle vertex in the singlet (s) and triplet (t) spin combination. The transversal particle-hole channel follows from $\Gamma_{ph}$ by (crossing) symmetry. Bottom: Fully irreducible vertex $\Lambda_{m(d)}$ for the two spin combinations. For all figures, the bare interaction $U$ has been subtracted from the vertices (reproduced from [26]).

Fig. 10 shows the reducible vertex $F$, the irreducible vertex in the particle-hole channel $\Gamma_{ph}$ and the fully irreducible vertex $\Lambda$ for the three-dimensional Hubbard model on a simple cubic lattice. For more details on the calculation of $\Lambda_{loc}$, see [26] and [22]. Also note that $\Gamma_{rloc}$ diverges at an interaction strength $U$ below the Mott-Hubbard metal-insulator transition, which signals the breakdown of perturbation theory [27].

2.2.1 Self consistency

After calculating $\Lambda_{loc}$, we can calculate the full vertex $F$ through the parquet equations, Fig. 7 and through the Schwinger-Dyson equation, Fig. 8 the non-local self energy $\Sigma$ and Green function $G$, as discussed in Section 2.1. Hitherto, all DΓA calculations have stopped at this point. That is, $F$ and $G$ are determined self-consistently but $\Lambda_{loc}$ is not recalculated.

However, in principle, one can self-consistency iterate the approach. From the new $G$ we can calculate a new $G_{loc}$. From this and $U$ we obtain a new vertex etc. until convergence, see Fig. 11. This self-consistency cycle is similar as that of the DMFT but now includes self-consistency on the two-particle ($\Lambda$) level.
Fig. 11: Flow diagram of the $D\Gamma A$ approach. Starting from a test Green function $G$ (e.g., that of DMFT), the local susceptibility and local full vertex $F_{\text{loc}}$ is calculated by solving an Anderson impurity model. To this end, the non-local Green function $G_0$ of an Anderson impurity is adjusted by Eq. (12) until this $G_0$ impurity model has the given interacting $G$. From $F_{\text{loc}}$ in turn, the inversion of the parquet and Bethe-Salpeter equations allow the calculation of the local fully irreducible vertex $\Lambda_{\text{loc}}$. This is the input of the parquet equations for calculating the non-local vertex $F$ and through the equation of motion and the Dyson equation, the $D\Gamma A$ self energy $\Sigma$ and Green function $G$. In a self-consistent calculation a new local vertex is calculated from $G$ etc. until convergence.

Please note that the Anderson impurity model has now to be calculated with the interacting $G_{\text{loc}}$ from $D\Gamma A$ which is different from the $G_{\text{loc}}$ of DMFT. As numerical approaches solve the Anderson impurity model for a given non-interacting Green function $G_0$, we need to adjust this $G_0$ until the Anderson impurity model’s Green function $G$ agrees with the $D\Gamma A$ $G_{\text{loc}}$. This is possible by iterating $G_0$ as follows:

$$[G_0^{\text{new}}(\nu)]^{-1} = [G_0^{\text{old}}(\nu)]^{-1} + [G_{\text{loc}}(\nu)]^{-1} - [G_{\text{loc}}^{\text{old}}(\nu)]^{-1},$$ (12)

until convergence. Here, $G_0^{\text{old}}$ and $G_{\text{loc}}^{\text{old}}$ denote the non-interacting and interacting Green function of the Anderson impurity model from the previous iteration. This $G_0$-adjustment is indicated in Fig. 11 by the secondary cycle.

Fig. 12 compares the $D\Gamma A$ self energies calculated this way, i.e., the $D\Gamma A$ full parquet solution, with DMFT and the exact solution. The results are for a simple one-dimensional Hubbard model with nearest neighbor hopping, six sites and periodic boundary conditions so that the exact solution is still possible by an exact diagonalization of the Hamiltonian. This can be considered as a simple model for a benzene molecule. The $D\Gamma A$ results have been obtained in a
Fig. 12: Self-energy $\Sigma_{ii+1}$ between neighboring sites on a six site Hubbard ring (see inset), comparing DMFT (zero non-local $\Sigma_{ii+1}$), $\Gamma_A$ and the exact solution. Parameters: $U = 2t$, $T = 0.1t$ with $t$ being the nearest neighbor hopping on the ring (reproduced from [30]).

“one-shot” calculation with the DMFT $G$ as a starting point. The good agreement between $\Gamma_A$ and the exact solution shows that $\Gamma_A$ can be employed for quantum chemistry calculations of correlations in molecules, at least if there is a gap [HOMO-LUMO gap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO)]. For molecules with degenerate ground states (i.e., a peak in the spectral function at the Fermi level), the agreement is somewhat less impressive; note that one dimension is the worst possible case for $\Gamma_A$.

In Fig. 12, the first parquet $\Gamma_A$ results have been shown. However, most $\Gamma_A$ calculations hitherto employed a simplified scheme based on ladder diagrams, see Fig. 13. These calculations neglect one of the three channels in the parquet equations: the particle-particle channel. Both the particle-hole ($ph$) and the transversal particle-hole channel ($ph$) are taken into account. These channels decouple for the spin and charge vertex and can hence be calculated by solving the simpler Bethe-Salpeter equations instead of the full parquet equations. If one neglects non-local contributions in one of the channels, it is better to restrict oneself to non-self consistent calculations since part of the neglected diagrams cancel with diagrams generated by the self consistency. Instead one better does a “one shot” calculation mimicking the self-consistency by a so-called $\lambda$ correction, see [29] for details. Physically, neglecting the particle-particle channel is justified if non-local fluctuations of particle-particle type are not relevant. Whether this is the case or not depends on the model and parameter range studied. Such particle-particle fluctuations are e.g. relevant in the vicinity of superconducting instabilities, where they need to be considered. In the vicinity of antiferromagnetic order on the other hand, the two particle-hole channels are the relevant ones (and only their magnetic spin combination). These describe antiferromagnetic fluctuations, coined paramagnons in the paramagnetic phase above the antiferromagnetic transition temperature. The $\Gamma_A$ results in the next section are for the half-filled Hubbard model. Here we are not only away from any superconducting instability, but at
3 Two highlights

3.1 Critical exponents of the Hubbard model

The Hubbard model is the prototypical model for strong electronic correlations. At half filling it shows an antiferromagnetic ordering at low enough temperatures $T$ – for all interaction strengths $U > 0$ if the lattice has perfect nesting. Fig. 14 shows the phase diagram of the half-filled three-dimensional Hubbard model (all energies are in units of $D = 2\sqrt{6}t \equiv 1$, which has the same standard deviation as a Bethe lattice with half bandwidth $D$). At weak interaction strength, we have a Slater antiferromagnet which can be described by Hartree-Fock theory yielding an exponential increase of the Néel temperature with $U$. At strong interactions, we have preformed spins with a Heisenberg interaction $J = 4t^2/U$ yielding a Heisenberg antifer-
Fig. 14: Phase diagram of the Hubbard model on a cubic lattice with nearest neighbor hopping $2\sqrt{6}t \equiv 1$. The dashed black lines show the DMFT Néel temperature $T_N$ and the DMFT crossover region from a paramagnetic metal (PM) to a paramagnetic insulator (PI). Non-local correlations reduce $T_N$ with good agreement between DΓA, DCA, lattice quantum Monte Carlo (QMC), as well as determinantal diagrammatic Monte Carlo (DDMC) before and after our DΓA results. Note, for the lowest $U$ value, Ref. [36] could only give an upper bound for $T_N$ which according to DDMC could be much smaller as indicated by the arrow.

romagnet with $T_N \sim J$. In-between these limits, $T_N$ is maximal.

All of this can be described by DMFT. However, since DMFT is mean-field w.r.t. the spatial dimensions, it overestimates $T_N$. This can be overcome by including non-local correlations, i.e., spatial (here antiferromagnetic) fluctuations. These reduce $T_N$. In this respect, there is a good agreement between DΓA, DCA and lattice QMC, see Fig. 14. The biggest deviations are observed for the smaller interaction strength. In principle, these differences might originate from the fact that the DΓA calculations are not yet self-consistent and only use the Bethe-Salpeter Eqs. (6,7) in the two particle-hole channels instead of the full parquet equations (5,10).

On the other side, however, we observe that long-range correlations are particularly important at weak coupling, cf. Section 3.2. Such long-ranged correlations cannot be captured by the cluster extensions of DMFT or lattice QMC since these are restricted to maximally $\sim 10$ sites in all three directions. More recent and accurate DDMC calculations on larger clusters [36] indeed show a smaller $T_N$ and, in contrast to DCA and lattice QMC better agree with DΓA, see Fig. 14. Even more important are long-range correlations in the immediate vicinity of the phase transition and for calculating the critical exponents. Each finite cluster will eventually show a mean-field exponent. In this respect, we could calculate for the first time the critical exponents of the Hubbard model [31]. In Fig. 15 we show the antiferromagnetic spin susceptibility which shows a mean-field-like behavior $\chi_{AF} \sim (T - T_N)^{-1}$ at high temperature (and in DMFT). In the vicinity of the phase transition however, long-range correlations become important and yield another
critical exponent $\chi_{AF} \sim (T - T_N)^{-2\nu}$ which agrees with that of the three dimensional Heisenberg model [32] (as is to be expected from universality). In contrast for the Falikov-Kimball model, the critical exponents calculated by the related dual Fermion approach [37] agree with those of the Ising model. Except for the immediate vicinity of the phase transition DMFT is nonetheless yielding a reliable description of the paramagnetic phase. At least this is true for one-particle quantities such as the self energy and spectral function, while the susceptibility [31] and entropy [35] show deviations in a larger $T$-interval above $T_N$.

### 3.2 Fate of the false Mott-Hubbard transition in two dimensions

As a second example, we recapitulate results for the interplay between antiferromagnetic fluctuations and the Mott-Hubbard transition in two dimensions. Even though the Hubbard model has been studied for 50 years, astonishingly little is known exactly. In one dimension it can be solved by the Bethe ansatz, and there is no Mott-Hubbard transition for the half-filled Hubbard model: For any finite interaction it is insulating. In infinite dimensions, on the other hand DMFT provides for an exact (numerical) solution. It has been one of the big achievements of DMFT to clarify the nature of the Mott-Hubbard transition which is of first order at a finite interaction strength [3, 4], see Fig. [16]. From cluster extensions of DMFT it has been concluded that the Mott-Hubbard transition is actually at somewhat smaller $U$ values and the coexistence region where two solutions can be stabilized is smaller, see Fig. [16]. However, again these cluster extensions are restricted to short-range correlations. In particular at low temper-
Fig. 16: Temperature $T$ vs. interaction $U$ phase diagram of the two dimensional Hubbard model on a square lattice with nearest neighbor hopping (all energies are in units of $D = 4t \equiv 1$, yielding the same standard deviation as for the 3D phase diagram). From local correlations (DMFT [39]) via short-range correlations (CDMFT [40] and variational cluster approximation (VCA) [38]) to long-range correlations ($D\Gamma A$ [31] and BSS-QMC [38]) the critical interaction strength for the metal-insulator transition is reduced to $U_c = 0$. The light gray lines denote the DMFT [33] and $D\Gamma A$ [31] $T_N$ (reproduced from [38]).

At reduced temperatures, there are strong long-range antiferromagnetic spin fluctuations which for example at $U = 0.5$ and $T = 0.01$ exceed 300 lattice sites [38]. The physical origin are antiferromagnetic fluctuations emerging above the antiferromagnetically ordered phase. In two dimensions this antiferromagnetic phase is restricted to $T = 0$ due to the Memin-Wagner theorem, but antiferromagnetic fluctuations remain strong even beyond the immediate vicinity of the ordered phase (at $T = 0$). These long-range antiferromagnetic spin fluctuations (paramagnons) give rise to pseudogap physics, where first only part of the Fermi surface becomes gaped but at lower temperatures the entire Fermi surface is gaped so that we have an insulating phase for any $U > 0$. Fig. 16 shows the development from local correlations (which only yield an insulating phase only relatively large $U$’s) to additional short-range correlations (which reduce the critical $U_c$ for the Mott-Hubbard transition) to long-range correlations (which reduce $U_c$ to zero). At large $U$, we have localized spins which can be described by a spin model or the Mott insulating phase of DMFT. At smaller $U$ we also have an insulator caused by antiferromagnetic spin fluctuations. These smoothly go over into the $T = 0$ antiferromagnetic phase, which is of Slater-type for small $U$. Since the correlation are exceedingly long-ranged, the nature of the low-temperature gap is the same as the Slater antiferromagnetic gap, even though there is no true antiferromagnetic order yet.
4 Conclusion and outlook

In the last years we have seen the emergence of diagrammatic extensions of DMFT. All these approaches have in common that they calculate a local vertex and construct diagrammatically non-local correlations from this vertex. In regions of the phase diagram where non-local correlations are short-range, results are similar as for cluster extensions of DMFT. However, the diagrammatic extensions also offer the opportunity to include long-range correlations on an equal footing. This allowed us to study critical phenomena and to resign the Mott-Hubbard transition in the two-dimensional Hubbard model to its fate (there is no Mott-Hubbard transition).

These were just the first steps. Indeed the diagrammatic extensions offer a new opportunity to address the hard problems of solid state physics, from superconductivity and quantum criticality to quantum phenomena in nano- and heterostructures. Besides a better physical understanding by hands of model systems, also realistic materials calculations are possible – by AbinitioDΓA. Taking the bare Coulomb interaction and all local vertex corrections as a starting point, AbinitioDΓA includes DMFT, GW and non-local correlations beyond within a common underlying framework. Both on the model level and for realistic materials calculations, there is plenty of physics to explore.

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