From infinite to two dimensions through the functional renormalization group

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We present a novel scheme for an unbiased, non-perturbative treatment of strongly correlated fermions. The proposed approach combines two of the most successful many-body methods, the dynamical mean field theory (DMFT) and the functional renormalization group (fRG). Physically, this allows for a systematic inclusion of non-local correlations via the fRG flow equations, after the local correlations are taken into account non-perturbatively by the DMFT. To demonstrate the feasibility of the approach, we present numerical results for the two-dimensional Hubbard model at half-filling.

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Introduction. – Correlated electron systems display undoubtedly some of the most fascinating phenomena of condensed matter physics such as high-temperature superconductivity and quantum criticality; and with the tremendous progress to cool and control atomic gases new many-body physics is explored nowadays. These systems pose a particular challenge for theory. In this paper, we discuss a new route for the theoretical treatment of strong correlations, which combines the strengths of two of the most successful approaches developed hitherto: dynamical mean field theory (DMFT) [1, 2] and functional renormalization group (fRG) [3–6].

DMFT represents the “quantum” extension of the classical (static) mean-field theory [2]. More formally, it provides the exact solution of a quantum many-body Hamiltonian in the limit of infinite spatial dimensions \((d \to \infty)\) [4]. DMFT allows hence for an accurate (and non-perturbative) treatment of the local part of the correlations. Among others, it provides the essential ingredients to describe the Mott-Hubbard metal-to-insulator transition in three-dimensional bulk systems [7, 8]. At the same time, the mean-field nature with respect to the spatial degrees of freedom implies that all non-local spatial correlations are completely neglected in DMFT.

A powerful technique to treat such non-local correlations is, instead, provided by the fRG. Its starting point is an exact functional flow equation [5], which yields the gradual evolution from a simple initial action to the full final action, that is, the generating functional of all one-particle irreducible vertex functions. The flow parameter (RG scale) is usually a momentum or energy cutoff. Expanding the functional flow equation yields an exact but infinite hierarchy of flow equations for the \(n\)-particle vertex functions, which for most calculations is truncated at the two-particle level. There have been many applications of such weak-coupling truncations to low-dimensional fermion systems with competing instabilities and non-Fermi liquid behavior (for a review, see [4]).

The approach we present here is coined DMF\(^2\)RG as the DMFT solution serves as a starting point of the fRG flow. DMF\(^2\)RG aims at overcoming the main restrictions of the two methods, i.e., the lack of non-local correlations in DMFT and the weak-coupling limitation in practical implementations of the fRG. The basic idea of the DMF\(^2\)RG is the following: We apply the fRG not starting from a problem without (or with trivial) correlations, but from a converged DMFT solution of the correlated system. This way, the local but possibly strong DMFT correlations, essential to capture the Mott-Hubbard physics, are fully taken into account from the very beginning. Non-local correlations beyond DMFT, particularly important for low-dimensional systems, will be systematically generated by the fRG flow. We note that alternative strong coupling starting points for the fRG flow were recently discussed for the Bose-Hubbard [10] and the single-impurity Anderson model [11].

Before turning to the DMF\(^2\)RG algorithm, let us mention alternative approaches proposed in the past to include non-local correlations beyond DMFT. They can be classified into cluster [12, 13] and diagrammatic extensions [14–19] of DMFT. The former ones are evidently complementary in nature to DMF\(^2\)RG, as they provide short-range correlation beyond DMFT, but at a high numerical cost, which poses significant limits to multiband calculations. Similarly as the diagrammatic extensions of DMFT, the DMF\(^2\)RG includes short- and long-range correlations on equal footing and improves the scaling with the number of orbitals. At the same time, instead of a simple selection of diagrams (e.g. second order perturbation theory, ladder, etc.), DMF\(^2\)RG exploits the more powerful RG and generates parquet-like diagrammatic corrections to DMFT. This way, DMF\(^2\)RG provides a systematic and unbiased treatment of electronic correlations beyond DMFT in all channels. Topologically the
same diagrams albeit with different Green’s functions and vertices are obtained in the proposed parquet implementations of DΓA [14] and multi-scale methods [18, 19]. This is however computationally much more demanding, and suffers from divergences of the two-particle irreducible vertex [14, 21].

Method. – A rather flexible and effective formulation of DMF²RG (see also the Supplementary Material section for further details) is obtained starting from the local (or “impurity”) action of DMFT

\[ S_{\text{DMFT}} = -\int_0^\beta d\tau d\tau' \sum_{i\sigma} \bar{c}_i(\tau) G^0_{\text{AIM}}(\tau - \tau')^{-1} c_i(\tau') + S_{\text{int}}. \]

Here, \( \bar{c}_i(\tau) \) are the Grassmann variables corresponding to the creation (annihilation) of a fermion with spin orientation \( \sigma = \uparrow, \downarrow \) on site \( i \), \( G^0_{\text{AIM}}(\tau - \tau') \) is the electronic-bath Green’s function of the auxiliary effective Anderson impurity model (AIM), which in a first step needs to be determined self-consistently in DMFT [7] (see left-hand side of Fig. 1), and \( S_{\text{int}} \) is a local interaction.

With this DMFT solution as a starting point, the fRG generates a flow to the finite-dimensional action of interest

\[ S_{\text{latt}} = -\int_0^\beta d\tau d\tau' \sum_{k\sigma} \bar{c}_{k\sigma}(\tau) G^0_{\text{latt}}(k, \tau - \tau')^{-1} c_{k\sigma}(\tau') + S_{\text{int}}, \]

where \( G^0_{\text{latt}}(k, \tau - \tau') \) is the free propagator of the finite dimensional system, which reads \( G^0_{\text{latt}}(k, \omega) = (i\omega - \epsilon_k + \mu)^{-1} \) in terms of Matsubara frequencies, the energy-momentum dispersion \( \epsilon_k \) and the chemical potential \( \mu \). In Fig. 1 the specific case of a 2D square lattice is shown.

For the DMF²RG scheme we now introduce a flow parameter \( \Lambda \) [28] so that \( G^0_{\text{latt}}(k, \omega) = (\omega - \epsilon_k + \mu)^{-1} \) in terms of Matsubara frequencies, the energy-momentum dispersion \( \epsilon_k \) and the chemical potential \( \mu \). In Fig. 1 the specific case of a 2D square lattice is shown.

The flow of DMF²RG hence gradually switches off the DMFT-bath and switches on the 2D hopping, including non-local correlations beyond DMFT. Neglecting three (and more) particle vertices, the flow equations [8, 24] for the self-energy and the two-particle vertex are shown in Fig. 1. The truncation of the hierarchy at the level of the two-particle vertex \( \Gamma \) relies on the assumption that the relevant physics is captured by the structure appearing on the two-particle level. Let us emphasize, however, that these (and more-) particle vertices are included on the local level by DMFT. This flow scheme results in the following single-scale propagator (defined as \( \partial G_\Lambda / \partial \Lambda |_{\Sigma^A=\text{fixed}} \))

\[ S_\Lambda(k, \omega) = G^0_\Lambda(k, \omega) \left[ G^0_{\text{latt}}(k, \omega)^{-1} - G^0_{\text{AIM}}(\omega)^{-1} \right] \] (3)

which includes the full Green’s function \( G_\Lambda(k, \omega) = \left[ G^0_\Lambda(k, \omega)^{-1} - \Sigma^A(k, \omega) \right]^{-1} \).
of the two-particle vertex function, i.e., in our case, $\Gamma$ in the particle-hole crossed channel, for zero transfer frequency ($\nu_2 - \nu'_1 = 0$), antiferromagnetic momentum transfer ($\mathbf{k}_2 - \mathbf{k}_1 = (\pi, \pi)$) and $\mathbf{k}_1 = (0, \pi)$, $\mathbf{k}_2 = (\pi, 0)$ computed by fRG, with interaction cutoff $\Lambda_{\text{int}}$ \cite{23} (inset) and DMF$^2$RG (main panel) for the two-dimensional half-filled Hubbard model at $U = 1$, at different (inverse) temperatures.

The Hubbard Hamiltonian reads \cite{34}

$$H = -t \sum_{\langle ij \rangle \sigma} c_{i \sigma}^\dagger c_{j \sigma} + U \sum_i n_{i \uparrow} n_{i \downarrow},$$

where $t$ denotes the nearest-neighbor hopping amplitude on a square lattice and $U$ the local Coulomb repulsion. In the following, we will define our energies in terms of $4t \equiv 1$, and fix the average particle density to half filling $n = 1$. In this case, the momentum transfer of $(\pi, \pm \pi)$ corresponds to perfect (antiferromagnetic) nesting on the square shaped Fermi surface.

We use a channel decomposition of the interaction vertex \cite{34, 35} with Matsubara frequency dependence of the self-energy and the interaction vertex. The momentum-dependence is taken into account by discretizing the Brillouin zone into patches with constant coupling function. If fine enough, this discretization captures the angular variation of the coupling function along the Fermi surface with good precision. For simplicity, we restrict ourselves to 8 patches, which already includes important physical aspects of the 2D system \cite{37}.

**Numerical results.** – Our calculations for the two-particle vertex function and self-energy are reported in Figs. 2 and 3 respectively. In Fig. 2 we plot the largest component ($g_{\text{max}}$) of the vertex function, which – at half-filling – is found in the particle-hole crossed channel for zero frequency and antiferromagnetic momentum transfer ($\pi, \pi$). The data, which refer to a weak-intermediate regime ($U = 1$), clearly show that the DMF$^2$RG mitigates the fRG tendency to a low-$T$ divergence of the flow: We still obtain a converged DMF$^2$RG result for $g_{\text{max}}$ at $\beta = 1/T = 10$, whereas the fRG flow for the vertex is manifestly divergent \cite{38}. Quantitatively, by fixing an upper-bound for $g_{\text{max}}$, we observe that the temperature at which it is reached is slightly decreased in DMF$^2$RG compared to fRG for moderate values of the interaction (up to $U = 0.75$) while is significantly decreased from $T \sim 0.125$ (fRG) to $\sim 0.085$ (DMF$^2$RG) at $U = 1$. This is attributed to the damping effect of the local correlations, included from the very beginning in the flow of DMF$^2$RG. We emphasize that this “divergence” is not associated with a true onset of a long-range order. In fact, fRG-schemes can be adapted to access also the disordered phase at lower $T$ \cite{34}, though such an extension goes beyond the scope of this work.

We now turn to the analysis of the self-energy results obtained with the DMF$^2$RG flow at the lowest temperature considered, i.e., $\beta = 10$. Here, the fRG flow diverges, and it is worth to compare the DMF$^2$RG results with the original DMFT data, see Fig. 3. As expected in 2D, the non-local correlations captured by the DMF$^2$RG strongly modify the DMFT (k-independent) results, determining a significant momentum dependence of the self-energy at low frequencies: While in DMFT a metallic solution, with a moderate Fermi-liquid renormalization of the quasi-particle mass, is obtained, in DMF$^2$RG we observe a strong enhancement of the imaginary part of the self-energy at the Fermi surface. In fact, at the “antinodal” point $(\pi, 0)$, where the largest value of $-\text{Im} \Sigma$ is found, the low-frequency behavior is mani-

**FIG. 2:** (Color online) Flow of the largest component ($g_{\text{max}}$) of the two-particle vertex function, i.e., in our case, $\Gamma$ in the particle-hole crossed channel, for zero transfer frequency ($\nu_2 - \nu'_1 = 0$), antiferromagnetic momentum transfer ($\mathbf{k}_2 - \mathbf{k}_1 = (\pi, \pi)$) and $\mathbf{k}_1 = (0, \pi)$, $\mathbf{k}_2 = (\pi, 0)$ computed by fRG, with interaction cutoff $\Lambda_{\text{int}}$ \cite{23} (inset) and DMF$^2$RG (main panel) for the two-dimensional half-filled Hubbard model at $U = 1$, at different (inverse) temperatures.

**FIG. 3:** (Color online) Comparison of the results for the imaginary part of the fermionic self-energy of the two-dimensional Hubbard model for $U = 1$, and $\beta = 10$, calculated within DMFT (k-independent, in black) and DMF$^2$RG, for different k-vectors (the color coding of the different $k$ is defined in the inset, note that the values of Im$\Sigma(k, \omega_n)$ for $k = (0, 0)$ and $(\pi, \pi)$ coincide because of the particle-hole symmetry). **Upper inset:** Scheme of the 8-patches discretization used for the calculations. **Lower inset:** $T^{-}$-dependence of the momentum-resolved spin correlation function $S(q, i\Omega = 0)$. 

\[ \beta = 6.0 \quad \beta = 8.5 \quad \beta = 10.0 \]
festly non quasi-particle-like, indicating the destruction of the Fermi surface in this region of the Brillouin zone. The trend of large non-local corrections to DMFT at the antinodal momentum and towards a pseudogap formation is similar to cluster-DMFT results [12, 10]. Deviations from the DMFT metallic results, albeit less marked, are found at the “nodal” point \((\frac{\pi}{2}, 0)\), for which one cannot exclude, at this temperature, a residual presence of strongly damped quasi-particle excitations. The significant reduction of \(-\text{Im}\Sigma\) w.r.t. DMFT, observed at \((0,0)\) or \((\pi,\pi)\), does not imply metallicity since these points are far away from the Fermi surface; and the real part of the self-energy (not shown) is also strongly enhanced w.r.t. DMFT. A further insight on the non-local correlations captured by the DMF\(^2\)RG is given by the analysis of the momentum/frequency-dependent susceptibilities, which in DMF\(^2\)RG can be extracted from the two-particle vertex. In the lower inset of Fig. 3 we show the DMF\(^2\)RG results for the momentum-resolved spin-susceptibility at zero frequency \(S(q,\Omega = 0)\). This quantity is most important at half-filling, where magnetic fluctuations predominate, and it is experimentally accessible, e.g., via neutron spectroscopy. Our results are in qualitative agreement with the QMC data of Refs. [11, 12] and show the major role played by antiferromagnetic fluctuations, with a pronounced peak at \((\pi,\pi)\), growing upon decreasing \(T\). The ferromagnetic fluctuations also get enhanced due to the van Hove singularity at the Fermi level.

In Fig. 4, we compare the DMF\(^2\)RG self-energy data with the \(\text{fRG}\). The comparison can only be performed at weaker coupling and/or higher \(T\) than in Fig. 3 as the \(\text{fRG}\) flow needs to converge. Our numerical data of Fig. 3 indicate that in the considered parameter region (same \(T\), but weaker interaction than in Fig. 3) the \(\text{fRG}\) and DMF\(^2\)RG yield qualitatively similar results for the \(k\) dependent self-energy. Considering that in DMF\(^2\)RG local correlations have been included non-perturbatively via DMFT, this confirms the validity of previous \(\text{fRG}\) analysis of the Hubbard model at weak and moderate interaction. At the same time, the applicability of DMF\(^2\)RG goes beyond the weak-to-intermediate coupling of the \(\text{fRG}\), allowing for the study of parameter regions where the Mott-Hubbard physics “already” captured by DMFT becomes important. Technically, a full treatment of this regime requires an improvement of the frequency parametrization of the 1PI vertex in the \(\text{fRG}\)-flow beyond the current frequency decomposition [35].

**Summary and outlook.** – We introduced the DMF\(^2\)RG approach, which exploits the synergy of local DMFT correlations and non-local correlations generated by the \(\text{fRG}\) flow. Applying DMF\(^2\)RG to the 2D Hubbard model, we find that, due to the inclusion of all local correlations by the DMFT starting point, the divergence of the flow for the interaction vertex is pushed to lower temperatures, where significant non-local corrections to DMFT are found. At the same time, in the temperature interval where both \(\text{fRG}\) and DMF\(^2\)RG converge, the self-energy results are qualitatively similar, supporting the results of previous \(\text{fRG}\) studies at weak-to-intermediate \(U\). Quantitatively, the most visible effect of DMF\(^2\)RG compared to \(\text{fRG}\) consists in a stronger \(k\)-dependence of the self energy for the considered parameters and a suppression of the “pseudocritical” temperature at which the vertex diverges. We emphasize, finally, the potential of the presented DMF\(^2\)RG approach to access the strong-coupling regime, where the Mott-Hubbard physics captured by DMFT will play a more important role and qualitative changes in the self-energy results are to be expected. The flexibility of the DMF\(^2\)RG scheme and its ability to avoid the sign-problem of a direct QMC treatment of non-local physics beyond DMFT look promising for future, unbiased studies of correlations in realistic multi-band models.

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SUPPLEMENTARY MATERIAL
From infinite to two dimensions through the functional renormalization group
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Derivation of the DMF²RG flow equations

Let us start by briefly recapitulating the standard fRG technique, which makes it easier to clarify how the DMFT algorithm can be combined with it. We consider an interacting problem

\[ S_{\text{latt}} = -\int_0^\beta d\tau d\tau' \sum_{k\sigma} c_{k\sigma}^\dagger(\tau) G_{\text{latt}}^0(k, \tau - \tau')^{-1} c_{k\sigma}(\tau') + S_{\text{int}}, \]

where \( G_{\text{latt}}^0 \) is the propagator of the Gaussian part, and all the terms beyond the Gaussian one are contained in \( S_{\text{int}} \); for the further notation see the paper. In general, the fRG procedure can be summarized conceptually in three steps:\[1,2]

1. First a "solvable" action \( (S_{\text{ini}}) \) is introduced as initial starting point. Here, "solvable" means that at the beginning the "problematic" parts of the original action are excluded (e.g., the degrees of freedom close to the Fermi level). Note that \( S_{\text{ini}} \) differs from the original \( S_{\text{latt}} \) only in its Gaussian part.

2. A one-parameter family of actions \( S^\Lambda \) is defined. These actions smoothly interpolate between the solvable action for the initial value of the parameter (i.e., if \( \Lambda = \Lambda_{\text{ini}} \), \( S^\Lambda_{\text{ini}} \equiv S_{\text{ini}} \)) and the physical one at the end (for \( \Lambda = \Lambda_{\text{fin}} \), \( S^\Lambda_{\text{fin}} \equiv S_{\text{latt}} \)). This corresponds to a continuous change of the Gaussian propagator from \( S_{\text{ini}} \) to \( S_{\text{latt}} \).

3. The evolution of all (1PI) \( m \)-particle vertex functions of the actions \( S^\Lambda \) as a function of \( \Lambda \) is determined from a set of coupled differential equations, called "flow equations".

The formal derivation of this procedure, as well as of the flow equations for the vertex functions is presented exhaustively in the literature, see, e.g., the recent reviews Refs. [1,2].

By integrating this set of differential equations, one can in principle evaluate exactly all 1PI \( m \)-particle vertex functions of the action \( S \) of the problem of interest by computing the flow from the corresponding vertex functions of the solvable model, independently on which specific choice was made for it. However, in the presence of a two-particle interaction, the hierarchy of flow equations couples the \( m \)-particle vertex function \( \Gamma^\Lambda_m \) with the \((m+1)\)-particle one, i.e., the set of flow equations is in general infinite. Hence, in practice one needs to truncate the equations: As an approximation, it is assumed that all the 1PI-vertex functions with \( m \) bigger than some value (typically \( m_{\text{max}} = 2 \)) are neglected. Within this approximate treatment, the choice of the initial action becomes obviously important.

More specifically, by retaining only the one-particle vertex function (self energy) and the two-particle vertex, and setting the three-particle vertex to zero, the truncated flow equations assume the form:

\[ \partial_\Lambda \Sigma^\Lambda = \Gamma^\Lambda_2 \circ S^\Lambda, \]
\[ \partial_\Lambda \Gamma^\Lambda_2 = \Gamma^\Lambda_2 \circ (S^\Lambda \circ G^\Lambda) \circ \Gamma^\Lambda_2. \]

Here the symbol "\( \circ \)" stands for the standard summation over all internal variables, i.e., momentum integration as well as spin and Matsubara frequency summation. At each vertex, energy, spin, and momentum is conserved according to the conventional diagrammatic rules. The symbols \( \Sigma^\Lambda, \Gamma^\Lambda_2, G^\Lambda \) and \( S^\Lambda \) stand respectively for the self energy, two-particle vertex, dressed Green’s function and single scale propagator, as defined in the main text. The initial condition for these differential equations, \( \Sigma_{\text{ini}}^\Lambda, \Gamma^\Lambda_2^\text{ini} \), are obtained by solving the initial “solvable” action \( S_{\text{ini}} \). We note, finally, that the Eqs. (2) and (3) correspond diagrammatically to the ones reported in Fig. 1 of the manuscript. Their explicit expression in terms of frequency, momenta and spin summations can be found, e.g., in Refs. [1 and 2].

Let us note that since the fRG flow-equations cannot be solved without a truncation (at a given loop level), the final results depend on the starting point. In this respect, any fRG flow is “biased towards” its starting point. The quality of the results depend on how well the fRG flow can build up the missing part of the physics.

The basic idea of the new DMF²RG scheme can be summarized as follows: differently from the conventional fRG approach, in DMF²RG we aim at including a major part of the correlated physics already at the level of the initial “solvable” action. This is certainly possible for the non-perturbative, but purely local, correlations of DMFT, because the DMFT solution of several models and realistic problems of solid state physics can be obtained both at the one and the two-particle level\[3-5\].

The formal implementation of this idea requires evidently to replace the initial action with a one describing the non-perturbative local physics of the DMFT solution and then to set up the flow to the final action \( S_{\text{latt}} \) of the desired
problem (where all correlations, namely also those beyond DMFT, are eventually included). Due to the flexibility of the fRG scheme, there are several ways to do this in practice. From a mathematical point of view, as DMFT corresponds to the exact solution of a quantum many body Hamiltonian in the limit of infinite dimensions \((d \rightarrow \infty)\), the most intuitive way might be realized by building up a “dimensional” flow from \(d = \infty\) to the actual dimensions (e.g., \(d = 2\) or \(3\)) of the problem of interest. In this case, one would start from the action of an infinite dimensional lattice (e.g., hypercubic) and the parameter \(\Lambda\) should gradually turn off the hopping in all directions, except the physical ones of the final problem.

This can be done considering the family of actions associated with the following Hamiltonians in the limit \(d \rightarrow \infty\):

\[
H^{\Lambda} = \sum_{k \sigma} \left\{ \frac{1}{\sqrt{2d}} \left[ f(\Lambda)\epsilon_{k_1k_2} + \Lambda\epsilon_{k_3...k_d} \right] - \mu \right\} c^\dagger_{k\sigma}c_{k\sigma} + U\sum_i n_{i\uparrow}n_{i\downarrow}.
\]

Here the momenta in the first sum are \(d\) dimensional: \(k = (k_1, k_2, ..., k_d)\) while the second sum extends over the lattice sites of a \(d\) dimensional lattice. The operators \(c^\dagger_{k\sigma}\) (\(c_{k\sigma}\)) create (annihilate) a fermion of momentum \(k\) and spin \(\sigma\), \(n_{i\sigma}\) is the number operator counting the fermions of spin \(\sigma\) at the lattice site \(i\).

To be specific, let us assume that the energies \(\epsilon_{k_1k_2}\) refer to a two dimensional square lattice with nearest neighbors hopping \(t\): \(\epsilon_{k_1k_2} = -2t(\cos k_1 + \cos k_2)\). The infinite dimensional limit of this lattice is obtained when \(\Lambda = 1\) and \(f(\Lambda) = 1\) taking \(\epsilon_{k_3...k_d} = -2t(\cos k_3 + ... + \cos k_d)\). The factor \(\frac{1}{\sqrt{2d}}\) accounts for the proper scaling: it guarantees that the kinetic energy does not diverge in the limit \(d \rightarrow \infty\). The terms \(\Lambda\) and \(f(\Lambda)\) are used to interpolate between the Hamiltonian in \(d\) and two dimensions. For example, assuming the following form for \(f(\Lambda)\):

\[
f(\Lambda) = 1 + (1 - \Lambda)(\sqrt{2d} - 1),
\]

one recovers the \(d \rightarrow \infty\) limit for \(\Lambda = 1\), while for \(\Lambda = 0\) one restores the two dimensional lattice Hamiltonian.

In spite of its intuitive picture, however, such “dimensional” flow equations, might be not the most suitable scheme to be adopted in practice. In fact, one should consider that in most of its applications, and in particular in those aiming at the realistic description of materials, DMFT is employed as an “approximation” for describing the local physics of a given finite-dimensional system, and no limit of infinite dimensions is actually taken. In fact, it would be rather cumbersome to define a rigorous and general procedure for connecting on a Hamiltonian level, case by case, the kinetic energy does not diverge in the limit \(d \rightarrow \infty\). The terms \(\Lambda\) and \(f(\Lambda)\) are used to interpolate between the Hamiltonian in \(d\) and two dimensions. For example, assuming the following form for \(f(\Lambda)\):

\[
f(\Lambda) = 1 + (1 - \Lambda)(\sqrt{2d} - 1),
\]

one recovers the \(d \rightarrow \infty\) limit for \(\Lambda = 1\), while for \(\Lambda = 0\) one restores the two dimensional lattice Hamiltonian.

As specified in the previous section, after defining the initial and the final actions, one must also set up a collection of one-parameter dependent actions smoothly interpolating between them. In the case of DMFT, a quite natural choice is a linear interpolation of the Gaussian part \((G^{\Lambda}_A(k, i\omega)^{-1})\) of the action from \(S^{\Lambda_{\text{ini}}} = S_{\text{DMFT}}\) (where \(S_{\text{DMFT}} = -\int_0^\beta d\tau d\tau' \sum_i \tilde{c}_{i\sigma}(\tau) G^{\text{A}_\text{AIM}}_{\text{A}_\text{AIM}}(\tau - \tau')^{-1} c_{i\sigma}(\tau') + S_{\text{int}}\)) to \(S^{\Lambda_{\text{fin}}} = S_{\text{latt}}\), which reads explicitly:

\[
G^{\Lambda}_A(k, i\omega)^{-1} = f(\Lambda)G^{\text{A}_\text{AIM}}_{\text{AIM}}(i\omega)^{-1} + [1 - f(\Lambda)]G^{\text{A}_\text{latt}}_{\text{latt}}(k, i\omega)^{-1},
\]
FIG. 1: (Color online) On the left, diagrammatic representation of the two-particle 1PI vertex function $\Gamma_2^\Lambda$; the arrows mark the position of the incoming and outgoing electrons. On the right, DMFT vertex function $\Gamma^{\Omega_1,\Omega_2,\Omega_3}$ as a function of $\Omega_1$ and $\Omega_3$ at fixed $\Omega_2$ ($\Omega_2 = 2\pi T n_1$; $n_2 = 20$) for the Hubbard model on a three dimensional cubic lattice with nearest neighbors hopping and $U = 0.5D$, $T = 0.038 D$ ($D$ being the half bandwidth). The color coded values are measured in units of $D$. Please note that in DMFT the main features of the vertex do not depend on the details of the lattice, but only on the bandwidth, therefore the following considerations apply in general. The white background color corresponds to the asymptotic value $U$ reached by the vertex. On the top of this, one can recognize three structures: i) a vertical line at $\Omega_1 = 0$, ii) a horizontal line at $\Omega_3 = 0$, and iii) a broader (hardly discernible) cross structure on the diagonals at $\Omega_1 = \pm \Omega_3$. The origin of the three structures has been analyzed in Ref. 4. While the structures i) and ii) are well described by the frequency dependence approximation described in the text, the cross structure is not captured by the approximation. Please notice that the white corners on the right of the density plot correspond to frequencies not included in the frequency window of our data set.

where $f(\Lambda)$ is an arbitrary smooth function of $\Lambda$ such that $f(\Lambda_{\text{ini}}) = 1$ and $f(\Lambda_{\text{fin}}) = 0$. For the sake of clarity, in the manuscript we have chosen $f(\Lambda) = \Lambda$ with $\Lambda_{\text{ini}} = 1$ and $\Lambda_{\text{fin}} = 0$, but, obviously, any alternative choice of $f(\Lambda)$ simply leads to an equivalent formulation of the truncated flow equations.

This choice of “cutoff” scheme $S^\Lambda$ according to Eq. (6) is similar to the “interaction cutoff” in the standard fRG, since it does not operate any selective cut on specific regions of the momentum and/or frequency space. The implementation of a frequency cutoff, may read

$$G^0_\Lambda(k, i\omega)\bigg|^{-1} = \theta(\Lambda - |\omega|)G^0_{\text{AIM}}(i\omega)\bigg|^{-1} + \theta(|\omega| - \Lambda)G^0_{\text{latt}}(i\omega, k)\bigg|^{-1},$$

where $\theta(x)$ is the Heavyside-step function. Evidently all possible cutoff schemes are equivalent in the case of a non-truncated flow. In the actual implementation however, a frequency- or momentum-cutoff, which can regularize infrared divergences of the problem, might be more suited, in particular, to study the regime in the proximity of (quantum) phase transitions. Its effective implementation, however, is numerically more involved than the simple cutoff of Eq. (6) and subject to future investigations.

**Parametrization of the two-particle vertex**

Here we give some details about the approximation employed on the frequency dependence of the 1PI two-particle vertex function for the single-band Hubbard model, studied in the manuscript. We use the conventions and definitions of Ref. 4 (in particular in “particle-hole notation”). Please note, however, that there the two-particle 1PI vertex is labeled $F$ and is momentum independent, while here it is called $\Gamma_2^\Lambda$ and can depend on the momentum.

In general for an SU(2) symmetric interaction and for a translationally invariant system the vertex function depends on two spins, three frequencies, and three momenta variables (see Fig. 1a):

$$\Gamma^{\Lambda, \nu\nu'\omega}(k_1', k_2'; k_1) := \Gamma_2^\Lambda\bigg(\nu k_1\sigma, (\nu' + \omega)k_2'\sigma' \bigg| \nu' k_1'\sigma', (\nu + \omega)k_2\sigma\bigg).$$

(8)
Here \( \nu \) and \( \nu' \) are fermionic Matsubara frequencies, while \( \omega \) is a bosonic Matsubara frequency. Physically this describes the scattering of a hole of energy \(-\nu\) with an electron of energy \(\nu + \omega\).

For the implementation of the frequency parametrization used in previous fRG studies\(^1\) it is however more convenient to adopt a frequency notation in terms of three bosonic Matsubara frequencies\(^1\) defined as follows:

\[
\begin{align*}
\Omega_1 &= \nu + \nu' + \omega, \\
\Omega_2 &= \nu - (\nu + \omega) = -\omega, \\
\Omega_3 &= \nu' + \omega - (\nu + \omega) = \nu' - \nu.
\end{align*}
\]

As for the spin indexes by exploiting the SU(2) symmetry we have

\[
\Gamma_{2,\uparrow\downarrow}^{A,\Omega_1,\Omega_2,\Omega_3}(k'_1, k'_2; k_1) = \Gamma_{2,\uparrow\downarrow}^{A,\Omega_1,\Omega_2,\Omega_3}(k'_1, k'_2; k_1) - \Gamma_{2,\uparrow\downarrow}^{A,\Omega_1,\Omega_2,\Omega_3}(k'_2, k'_1; k_1).
\]  

(12)

Hence, we can concentrate on the vertex \( \Gamma_2^{A,\Omega_1,\Omega_2,\Omega_3}(k'_1, k'_2; k_1) \) only (all the other spin combinations can be obtained by symmetry).

Even by restricting ourselves to the \( \uparrow\downarrow \) sector, the vertex function \( \Gamma_2^{A,\Omega_1,\Omega_2,\Omega_3} \) displays, in general, a rather complicated structure in momentum and frequency space. However, the efficiency of our first DMF\(^2\)RG calculations could be improved by parametrizing the frequency dependence of \( \Gamma_2^{A,\Omega_1,\Omega_2,\Omega_3} \) following the previous experience of an fRG study of the AIM\(^2\), where the following approximation for the frequency dependence of the vertex:

\[
\Gamma_{2,\uparrow\downarrow}^{A,\Omega_1,\Omega_2,\Omega_3}(k'_1, k'_2; k_1) \approx U + \tilde{\Gamma}_{2,pp}^{A,\Omega_1}(k'_1, k'_2; k_1) + \tilde{\Gamma}_{2,\phi-d}(k'_1, k'_2; k_1) + \tilde{\Gamma}_{2,\phi-c}(k'_1, k'_2; k_1),
\]  

(13)

is proposed.

This corresponds to approximating the complicated dependence of \( \Gamma_2^{A,\Omega_1,\Omega_2,\Omega_3} \) on the three bosonic frequencies \( \Omega_1, \Omega_2, \Omega_3 \), assuming that the scattering amplitude among two particles can be completely decomposed in three different channels (pp, pp – d, ph – c). This assumption is not exact for a generic \( U \), as one could immediately see already by looking at the DMFT vertex function, which represents the input for DMF\(^2\)RG. However, as described in detail in Refs.\(^1\)\(^1\) it is consistent with the lowest-order perturbation theory for \( \Gamma_2^{A,\Omega_1,\Omega_2,\Omega_3} \). Following Ref.\(^1\)\(^1\) one can derive the flow equations directly for the functions \( \Gamma_{2,\uparrow\downarrow}^{A,\Omega_1,\Omega_2,\Omega_3} \), \( \tilde{\Gamma}_{2,pp}^{A,\Omega_1} \), \( \tilde{\Gamma}_{2,\phi-d} \), \( \tilde{\Gamma}_{2,\phi-c} \). This is possible because one can associate each function with a specific channel: particle-particle, particle-hole direct and particle-hole crossed.

In fact, it has been shown that this approximation correctly describes the vertex structures up to \( \mathcal{O}(U^3) \) and it is expected to be reliable for moderate values of \( U \). On the other hand, increasing the \( U \) value, structures not captured by Eq. \(^1\)\(^1\), and arising from higher order diagrams (like the diagonal ones at \( \Omega_1 = \pm \Omega_3 \) in Fig.\(^1\)) will become more important, making the approximation unreliable. This is the reason why we only present results for moderate \( U \) values in the paper.

The only point left to discuss is how to extract the initial condition for the three functions in Eq. \(^1\)\(^1\) from the frequency dependent DMFT vertex \( \Gamma_{DMFT}^{\Omega_1,\Omega_2,\Omega_3} \) which contains more information than necessary. By looking at Fig.\(^1\), one sees that the problem consists in how to get rid of the cross structure (labeled \( \text{iii} \)) which depends on all the frequencies. However, the structure under consideration fades out becoming gradually broader and less intense as the third frequency is increased. Therefore to extract one of the three functions, say, e.g., \( \tilde{\Gamma}_{2,pp}^{A,\Omega_1} \), it suffices to take a cut in \( \Gamma_{DMFT} \) keeping \( \Omega_2 \) and \( \Omega_3 \) fixed at some very large values \( \Omega_2^3 \) and \( \Omega_3^3 \):

\[
\tilde{\Gamma}_{2,pp}^{A,\Omega_1}(k'_1, k'_2; k_1) = \Gamma_{DMFT}^{\Omega_1,\Omega_2^3,\Omega_3^3}.
\]  

(14)

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