Introduction. The discovery of the spin-triplet superconductivity (SC) inside the ferromagnetic (FM) phase of heavy-fermion compound UGe$_2$ sparked an intense discussion about the cause of such coexistence. Although the spin-triplet paired phase has been known to appear in the condensed $^3$He and most likely in Sr$_2$RuO$_4$, until its discovery in UGe$_2$ there was no convincing example for a strongly FM material hosting SC.

Specifically, the phase diagram for UGe$_2$ on the temperature-pressure (T–p) plane contains both SC and two FM phases, with stronger and weaker magnetization, usually referred to as FM2 and FM1, respectively, as well as paramagnetic phase (PM), with the phase transitions between them of the 1st order for low temperature, $T \lesssim 7K$. The FM–SC coexistence is strongly suggestive of a single mechanism based on magnetic correlations which is responsible for both FM and SC appearance and thus should be treated on equal footing, as e.g. in UGe$_2$ both phases disappear at the same pressure. Another indication of the coupled nature of both phases is that SC dome on the T–p plane coincides with the phase transition between FM2 and FM1. Thus, we address here in detail the question of microscopic origin of the observed ferromagnetism, as it should bring us closer to determining the mechanism of superconductivity. The question related to the inclusion of SC requires a separate study (see discussion at the end).

Experimental observations suggest that the ferromagnetism in UGe$_2$ has an itinerant nature and is mediated by the uranium 5f electrons. Delocalization of the 5f electrons can be interpreted as resulting from hybridization of 5f originally atomic states with those from conduction bands derived from p states due to Ge and d-s states due to U. This is supported by a noticeable difference of the effective paramagnetic moment per uranium atom in this compound with respect to the corresponding atomic value for either $f^3$ or $f^2$ configurations, as well as from a fractional value of the magnetization relative to the moment saturation. This means that the Hund’s rule coupling in the atomic sense is broken, and the itineracy of 5f electrons is the source of the band ferromagnetism in which Hund’s ferromagnetic interaction plays a role in combination with the stronger intra-atomic Coulomb interaction. This also means that the f-electron orbital degeneracy is not essential, but the role of the hybridization is.

Apart from other theories concerning origin of FM in considered class of materials, there exist a phenomenological rationalization of the magnetic properties within a rigid-band Stoner approach, which requires introduction of an ad hoc two-peaked structure of density of states (DOS) near the Fermi surface (FS). Our purpose is to invoke a microscopic description starting from the Anderson-lattice model (ALM) which is appropriately adapted to the heavy-fermion compound UGe$_2$. This comprises a relatively simple quasi-two-dimensional electronic structure. From such starting point an effective non-rigid two-band description arises naturally and allows for a detailed rationalization of magnetic and electronic properties, at least on a semiquantitative level. Additionally, as the correlations among 5f electrons are sizable, an emergence of the Stoner-like picture of FM can be accounted for only with inclusion specific features coming from the electronic correlations. Although the resulting rationalization of the physical properties is semiquantitative in nature, it provides in our view a coherent picture of a number of properties.

Model. We solve ALM by means of a variational treatment with the Gutzwiller wave function, $\Psi_G = \prod_i \tilde{P}_i |\psi_0\rangle$, where $\tilde{P}_i$ is the operator projecting out part of double occupancies from the uncorrelated ground-state $|\psi_0\rangle$ at site $i$. We have extended the standard approach to the statistically consistent form (SGA method). Explicitly, we start with the ALM Hamiltonian, with an applied magnetic field introduced via the Zeeman term ($h \equiv \frac{1}{2} g \mu_B H$), i.e.,

\[
\hat{H} = -\mu \hat{N} = \sum_{i,j,\sigma} t_{ij}^f c_{i,\sigma}^\dagger c_{j,\sigma}^\phantom{\dagger} - \sum_{i,\sigma} (\mu + \sigma h) \hat{n}_{i,\sigma}^c + \sum_{i,\sigma} (\epsilon_f - \mu - \sigma h) \hat{n}_{i,\sigma}^f + U \sum_i \hat{n}_{i,\uparrow}^f \hat{n}_{i,\downarrow}^f + V \sum_{i,\sigma} (\hat{f}_{i,\sigma}^\dagger \hat{c}_{i,\sigma}^\phantom{\dagger} + \text{h.c.}) \tag{1}
\]
where primed sum denotes summation over all lattice sites \( i \neq j \). \( f \) and \( \bar{c} \) are operators related to \( f \)- and \( c \)-orbitals respectively, with spin \( \sigma = \uparrow, \downarrow \). We have also defined the total number of electrons operator as \( \hat{N} \), and for the respective orbitals and spins as

\[
\hat{n}_{i,\sigma}^f = \hat{c}_{i,\sigma}^\dagger \hat{c}_{i,\sigma}, \\
\hat{n}_{i,\sigma}^d = \hat{\bar{c}}_{i,\sigma}^\dagger \hat{\bar{c}}_{i,\sigma}.
\]

In our model, we consider finite intra-\( f \)-orbital Coulomb interaction \( U \), the on-site inter-orbital hybridization \( V \), the hopping amplitude between the nearest-neighbor sites, and the atomic level for \( f \)-states placed at \( \epsilon_f = -3|t| \). In the following \(|t|\) is used as energy unit.

First, we would like to evaluate the ground-state energy, \( E_G \equiv \langle \psi_G | \hat{H} | \psi_G \rangle / \langle \psi_G | \psi_G \rangle \). Applying the usual procedure\(^{21,22}\), called Gutzwiller approximation (GA), we simplify the projection to the local sites on which the operators from (1) act. In that manner one obtains the effective single-particle Hamiltonian in a momentum space with renormalized hybridization by the Gutzwiller narrowing factor \( \frac{1}{2} \), namely

\[
\hat{H}_{\text{GGA}} \equiv \sum_{k,\sigma} \Psi^\dagger \left( \begin{array}{cc} \epsilon_k - \sigma \mu & \sqrt{\sigma} V \\ \sqrt{\sigma} V & \epsilon_k - \sigma \mu \end{array} \right) \Psi + AUd^2, \\
(2)
\]

where we have defined \( \Psi^\dagger \equiv (\hat{c}_{k,\sigma}^\dagger, \hat{\bar{c}}_{k,\sigma}^\dagger) \), \( \Lambda \) denotes number of lattice sites, and \( d^2 \) is the probability of having doubly occupied \( f \)-orbital that we optimize variationally. In order to ensure that variationally calculated polarization and the \( f \)-level occupancy would coincide with those coming from the self-consistent procedure\(^{23}\), we modify our effective Hamiltonian (2) by introducing additional constraints on the polarization \( (m_f) \) and the number \( \left( n_f \right) \) of \( f \)-electron states via the Lagrange-multiplier method. The effective Hamiltonian with the constraints takes now the form

\[
\hat{H}_{\text{GGA}} \equiv \hat{H}_{\text{GGA}} - \lambda_f^f \left( \sum_{k,\sigma} \hat{n}_{k,\sigma}^f - \Lambda n_f \right) - \lambda_m^f \left( \sum_{k,\sigma} \sigma \hat{n}_{k,\sigma}^f - \Lambda m_f \right) \\
= \sum_{k,\sigma} \Psi^\dagger \left( \begin{array}{cc} \epsilon_k - \sigma \mu & \sqrt{\sigma} V \\ \sqrt{\sigma} V & \epsilon_k - \sigma \mu \end{array} \right) \Psi + \Lambda (Ud^2 + \lambda_f^f n_f + \lambda_m^f m_f). \\
(3)
\]

Those constraint parameters \( \lambda_f^f \) and \( \lambda_m^f \) are also determined variationally. They play a role of nonlinear self-consistent fields acting on the charge and the spin degrees of freedom respectively. Diagonalization of (3) in this spatially homogeneous case leads to four branches of eigenenergies, \( E_{k\sigma}^\pm \), representing two spin-split hybridized bands \( E_{k\sigma}^\pm \). In order to determine the equilibrium properties of the system, we need to find the minimum of the generalized Landau grand-potential functional \( \mathcal{F} \),

\[
\frac{\mathcal{F}}{\Lambda} = -\frac{1}{\Lambda \beta} \sum_{k\sigma} \ln[1 + e^{-\beta E_{k\sigma}^\pm}] \\
+ (\lambda_f^f n_f + \lambda_m^f m_f + Ud^2), \\
(4)
\]

where \( b = \pm \). Effectively, it leads to the set of five nonlinear equations, \( \frac{\partial \mathcal{F}}{\partial \lambda_f^f} = 0 \) for \( \lambda_f^f \equiv \{ d, n_f, m_f, \lambda_f^f, \lambda_m^f \} \).

However, due to the fact that the total number of electrons remains constant when the pressure or magnetic field is applied, we need to satisfy equation for the chemical potential \( \mu \) via the condition

\[
\mu = \frac{1}{\Lambda} \sum_{k\sigma} \sigma f(E_{k\sigma}^b), \\
(5)
\]

with \( f \) being the Fermi distribution. The equilibrium thermodynamic potential functional defines also the ground state energy, \( E_G = \mathcal{F}_0 + \Lambda \mu n \), where subscript \( '0' \) denotes the optimal values. After carrying out the minimization, we can also calculate total spin polarization from

\[
m \equiv m_c + m_f = \frac{1}{\Lambda} \sum_{k\sigma} \sigma f(E_{k\sigma}^b). \\
(6)
\]

The numerical calculations with the precision of at least \( 10^{-7} \) were carried out for a two dimensional, square lattice of \( \Lambda = 512 \times 512 \) size, and for low temperatures \( \beta = 1/k_B T \geq 1500 \), emulating the \( T \rightarrow 0 \) limit.

**Results.** First, we analyze FM and PM solutions in the field absence. In Figure 1 we draw phase diagram on total filling–hybridization strength plane. For low hybridization, FM phases are favored due to the negative balance between increase of the kinetic and decrease of the Coulomb energies, caused by a relative shift of the spin-resolved DOS. This is visualized by the diminution of the spin-subband overlap up to the FS – cf. Figs. 1 and 7. The appearance of a spontaneous polarization, as a result of a competition between the kinetic and the Coulomb energies, is in fact the feature of the Stoner mechanism for the band FM onset. In comparison to the usual single-band (e.g. Hubbard) model, we can distinguish in a natural manner between the two FM phases. The first (FM1) appears when the chemical potential is placed in the hybridization gap, between the spin subbands of the lower hybridized band which is characterized also by the magnetization equal to \( m = 2 - n \) (cf. Fig. 1). In that situation, only the spin-minority carriers are present at and near FS. The second phase (FM2) emerges when we further lower the hybridization and thus the chemical potential enters the majority spin-subband DOS (cf. Fig. 1b), giving rise to a step (discontinuous) increase in magnetization (cf. Fig. 2b). In the limit of strong hybridization, for a fixed total filling, when the correlations weaken due to lowering of \( f \)-orbital average occupancy \( n_f \lesssim 0.85 \), c.f. Fig. 2b), the kinetic energy gain outbalances a subsequent reduction of the average Coulomb interaction and PM phase is energetically favorable. Similar mechanism for the formation of FM and in particular, characterization of phases, was studied before in Refs.\(^{23,26}\).

We presume that the main effect of the pressure exerted on the material can be modeled by a concomitant strengthening of the hybridization amplitude. Thus,
from Fig. 1 it can be seen that for the total filling \( n \) in the range 1.55–1.75, the sequence of phases and the order of the transitions are the same, as those found experimentally for UGe\(_2\) by increasing the pressure\(^{10,11}\). As a representative band filling we have selected \( n = 1.6 \) marked by the vertical line in Fig. 1. In fact, as we compare the magnetization versus hybridization along the traced line (cf. Fig. 1) with the corresponding experimental data\(^{4}\) (cf. Fig. 2) we find good qualitative resemblance. Moreover, the magnetization differentiation among the orbitals (cf. Fig. 2) is in agreement with the neutron scattering data\(^{10,11}\) at ambient pressure (in our model \(|V| \approx 0.5\), where it was found that almost exclusively electrons from uranium atoms (\(f\)-orbital) contribute to the ferromagnetism. In our Stoner-like picture it is resulting from the fact that the competition between Coulomb repulsion and hybridization-induced itineracy concerns mainly the \(f\) electrons. Furthermore, as for low hybridization (in FM2 phase) we obtain a small compensating polarization due to the \(c\)-electrons, we suggest that the experimentally observed small negative magnetization between the uranium atoms at ambient pressure\(^{11}\) may come from the delocalized cloud of conduction electrons.

Our microscopic description of the phase transitions induced by the change of the FS topology compares also favorably with the electronic-state features of UGe\(_2\) derived from de Haas van Alphen oscillations\(^{18,19}\). In Ref. 18 it is suggested, that the majority spin FS disappears in the FM1 phase, in complete accord with the character of DOS presented in the Fig. 1. We also reproduce the feature of an abrupt change of the FS at the FM1–PM phase transition\(^{18,19}\) (cf. Fig. 2). Namely, it corresponds here to the step change of the chemical potential position merging into both bands. Furthermore, in the experimental data at the metamagnetic phase transition there is observed significant enhancement of the quasi-particle mass renormalization\(^{19}\). As it is proportional to the DOS at Fermi level, in Fig. 2(b) we provide the corresponding behavior, which can be understood within our model by the chemical potential crossing high hybridization peak in the majority spin subband. The transition leads then to a step change of FS only in the majority spin subband, while the minority subband evolves rather continuously, what is also seen experimentally\(^{19}\).

For the sake of completeness, we have shown in the inset in Fig. 2 the pressure dependence of the \(T^2\) term.
of resistivity as it should have roughly the same dependence as squared DOS at the FS, versus $|V|$ (we assume that the Kadowaki-Woods scaling holds). However, the jump that we obtain at the FM1 - PM transition has not been observed in the resistivity measurements.

In the applied field, our model is also in good agreement with available experimental data for UGe$_2$. In Figure 3c, we display phase diagram on hybridization–applied-magnetic-field plane that corresponds to that determined experimentally (cf. Fig. 3b inset). Similarly as in Ref. [1], the magnetization at the phase transition between FM1 and FM2 triggered by the applied magnetic field, starts from the same baseline, independently of the hybridization strength (cf. Fig. 3b). However, one should note that, due to the fact that pressure changes not only the hybridization magnitude, but also other microscopic parameters, we are not able to reproduce the magnetization cascade with the increasing magnetic field when crossing the transitions.

The next feature found in UGe$_2$ at ambient pressure is an initial lack of measurable polarization on the germanium atoms with the increasing magnetic field, as inferred from the neutron scattering data. In our model we find a similar trend. For low hybridization ($|V| \approx 0.5$ emulating ambient pressure), c-electrons polarization increases slowly, and even up to $h \approx 0.1$ it is negligible (cf. Fig. 3).

Remarks. With the simple but powerful technique based on the generalized Gutzwiller ansatz (SGA method), applied to the Anderson lattice model, we have constructed a microscopic model of FM in UGe$_2$. Namely, we are able to reproduce main experimental features observed at low temperature, by applying either pressure or magnetic field (cf. Figs. 2a and 3a). FM properties can be rationalized within the simplest hybridized two-orbital model, without taking into account the f-orbital degeneracy, i.e., by effectively incorporating both the Coulomb and the Hund’s-rule interaction into an effective interaction $U$, as would be also the case in the Hartree-Fock approximation.

To determine the stability of SC inside FM phase, the present approach should be extended to account for the Hund’s-rule interaction explicitly what can be crucial for a formation of the unconventional triplet SC$^{16,27,28}$. If this is the case, it can be triggered even by a purely repulsive Coulomb interaction in conjunction with the residual Hund’s rule coupling, as discussed in Refs. 27 and 28. This issues require a separate analysis. Another path for discussing the coexistence of SC with FM could be going beyond the Gutzwiller approximation, where we account also for the more distant correlations when determining the effective Hamiltonian$^{29,30}$. Here, the central question is whether the spin triplet pairing should be treated on the same footing as ferromagnetism, i.e., appears already in direct space formulation$^{14,31–33}$ or is it mediated by collective spin fluctuations in ferromagnetic phase$^{14,31–33}$ among already well defined quasiparticles. A crossover from the latter to the former approach is expected to take place with the increasing strength of the repulsive Coulomb interaction $U$.

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