The last decade has seen a rapid development of methods at the two-particle (2P) processes. However, these are often performed with significant approximations or are limited to general definitions [1,2]. For this reason, corresponding analytical or numerical calculations correspond to a simple path of extracting its value directly from the lowest Matsubara frequency. We showcase its strength by applying it to the single impurity and the periodic Anderson model as well as to the Hubbard model. Our results represent a significant progress for the general understanding of quantum field theory at the two-particle level and allow for tracing the limits of the physics captured by perturbative approaches for correlated systems.

DOI: 10.1103/PhysRevLett.126.056403

Introduction.—The goal of any successful theory is to extract essential features of the phenomena of interest from the complexity of the physical world, neglecting all superfluous pieces of information. This objective is particularly crucial for the cutting-edge quantum field theory (QFT) approaches designed to describe many-electron systems in the presence of strong correlations.

Presently, one can rely on a solid textbook interpretation [1,2] of the QFT formalism describing the single-particle (1P) processes, measurable, e.g., by (angular resolved) direct and inverse photoemission [3] or scanning tunneling microscopy [4,5]. Crucial information about the metallic or insulating nature of a given many-electron problem, as well as quantitative information about the electronic mass renormalization $Z$ and quasiparticle lifetime $\tau$ is encoded in the momentum and energy dependence of the electronic self-energy $\Sigma$. If the temperature $T$ is low enough, even a quick glance at the low-energy behavior of $\Sigma$, either in real or in Matsubara frequencies, yields a qualitatively reliable estimate of the most important physical properties.

The situation is clearly different on the two-particle (2P) level, which can be experimentally accessed by, e.g., inelastic neutron scattering [6,7]. Because of the complex physical mechanisms at play, the related textbook knowledge is mostly limited to general definitions [1,2]. For this reason, corresponding analytical or numerical calculations are often performed with significant approximations or with a black-box treatment of the 2P processes. However, the last decade has seen a rapid development of methods at the forefront of the many-electron theory [8–10], for which generalized 2P correlation functions are the key ingredient. This is reflected in an increasing effort to develop the corresponding formal aspects and algorithmic procedures [8–31]. At the same time, the rather poor physical understanding of the 2P processes remains largely behind the requirements of the most advanced QFT methods. Interesting progress has been recently reported [32,33] on the relation of 1P Fermi-liquid parameters to 2P scattering functions. Ideally, however, one would like to be able to interpret the physics encoded at the 2P level with a similar degree of confidence as for the 1P processes.

In our Letter, we make a significant step forward in this direction: We identify the fingerprints of two major hallmarks of strong correlations in the generalized charge susceptibility. In particular, we pinpoint the frequency structures encoding the formation of local magnetic moments as well as of their Kondo screening. In this perspective, we also show how the Kondo temperature $T_K$ corresponds to a specific property of the generalized charge susceptibility, allowing for an alternative, simple path of extracting its value directly from the lowest Matsubara frequency data.

We recall that the Kondo problem [34] provides a paradigm for a variety of physical effects [35–39] involving strong electronic correlations. Local moment formation and Kondo screening are also a crucial ingredient of the physics described by the dynamical mean-field theory (DMFT) [40] through the solution of a self-consistently determined auxiliary Anderson impurity model (AIM).
Learning how to extract important physical information from the generalized susceptibility represents a substantial improvement for the understanding of quantum many-electron physics at the 2P level. Further, having this information at hand also enables us to draw conclusions on two relevant theoretical questions: (i) The relation of the recently reported multifaceted manifestations [41] of the breakdown of perturbation theory, such as the divergences of the irreducible vertex functions [15,42–49] and the crossing of multiple solutions [41,45,46,48,50–53] of the Luttinger-Ward functional, with the local moment physics and its Kondo screening; (ii) the built-in limits of advanced perturbative approaches to describe these fundamental physical effects.

How to read two-particle quantities.—We start from the definition of the generalized local susceptibility [10,12,54]

$$\tilde{\chi}^{\nu\nu}(\nu, \nu', \Omega) = G^{(2)}_{\nu\nu}(\nu, \nu', \Omega) - T^{-1}G(\nu)G(\nu')\delta_{\nu\nu}\delta_{\nu\nu'}$$

in terms of the 2P ($G^{(2)}$) and 1P ($G$) Green’s functions, where $\nu, \nu'$ and $\Omega$ are fermionic and bosonic Matsubara frequencies, respectively, and $\sigma, \sigma' = \{ \uparrow, \downarrow \}$ spin indices. As we show in the following for repulsive interactions, the generalized charge susceptibility $\tilde{\chi}^{\nu\nu}(\Omega) = \tilde{\chi}^{\uparrow\uparrow}(\Omega) + \tilde{\chi}^{\downarrow\downarrow}(\Omega)$ allows for the best readability of the underlying physics at the 2P level. Furthermore, the physical response of this sector captures the fundamental properties of any interacting electron system. We recall that the physical response function ($\chi$) is obtained from the generalized susceptibility $\tilde{\chi}^{\nu\nu}(\Omega)$ by summing over the fermionic Matsubara frequencies $\nu, \nu'$ [54]. The static charge response $\chi(\Omega = 0)$ reads

$$\chi = T^2 \sum_{\nu\nu'} \tilde{\chi}_{\nu\nu'} = T^2 \sum_{\nu\nu'} (\tilde{\chi}_{\uparrow\uparrow}^{\nu\nu} + \tilde{\chi}_{\downarrow\downarrow}^{\nu\nu}).$$

We start by analyzing the arguably simple case of an isolated atom with a repulsive interaction $U$ (Hubbard atom, HA), where analytic expressions are also available [12,48]. This represents the purest realization of local moment physics, which hence provides an ideal baseline for the interpretation of the more interesting cases discussed below. In Fig. 1 (upper panels), we show an intensity plot of $\tilde{\chi}^{\nu\nu}$ (normalized by $T^2$) for $U = 5.75$ [84], half filling (where $\tilde{\chi}^{\nu\nu}$ is real [12,48]) and different temperatures. At high temperature ($T_{\text{high}} = 2$, left panel), the overall frequency structure consists of a large positive-valued diagonal (yellow and red) and a weak negative cross structure (blue). This corresponds to a typical perturbative behavior [12,16], dominated by the diagonal bubble term $\tilde{\chi}^{\nu\nu}_0 = \tilde{\chi}_{0,\uparrow\uparrow}^{\nu\nu} = -\partial_{\nu\nu}G(\nu)^2/T$. Correlation effects are washed out for $T \gtrsim U$, consistent with the feasibility of high-$T$ expansions.

The situation changes radically when reducing $T$: in the intermediate ($T_{\text{int}} = 0.1$) and low ($T_{\text{low}} = 1/60 \approx 0.017$) temperature regime (central and right panel), one observes a strong damping of all diagonal elements of $\tilde{\chi}^{\nu\nu}$. The effect is more pronounced at low frequencies, as the sign of $\tilde{\chi}^{\nu\nu}$ becomes even negative (blush colors) for $|\nu| \lesssim U$ [48] (black square). This major feature is accompanied by the appearance of small positive off-diagonal elements (yellow). The net effect is a suppression of the physical susceptibility $\chi$, see Eq. (2), which occurs when the thermal energy is no longer large enough ($T \sim \nu < U$) to counter the formation of a local moment driven by $U$, eventually yielding an exponentially small $\chi \sim e^{-U/\Omega}$ for $T \to 0$. Altogether, the low-$T$ HA results illustrate how the onset of a pure local moment is encoded in the charge sector: a progressive emergence of a nonperturbative sign structure in $\tilde{\chi}^{\nu\nu}$, which is the opposite image of the perturbative one (left panel). This also induces several negative eigenvalues of $\tilde{\chi}^{\nu\nu}$, responsible for the breakdown of perturbative expansions [41].

Let us now examine how this picture changes when the HA system is connected to an electronic bath (here: with a flat DOS of bandwidth $W = 20$ and hybridization $V = 2 < U = 5.75$ [84]), corresponding to the well-known Anderson impurity model (AIM). By comparing the results of $T^2\tilde{\chi}^{\nu\nu}$ (central-row panels of Fig. 1, computed with w2dynamics [54]) to those of the HA, we observe almost no difference at $T_{\text{high}}$. This is not surprising as thermal fluctuations prevail over both correlation ($U$) and hybridization ($V$) effects in this case. Upon lowering $T$ to $T_{\text{int}}$, we enter the local moment regime of the AIM. This is reflected in a qualitatively similar evolution as seen in the HA: a progressive suppression of the diagonal entries of $\tilde{\chi}^{\nu\nu}$, turning negative in the low-energy sector (black square), accompanied by positive, yet smaller, off-diagonal contributions, with an overall freezing effect on the local density fluctuations [see Eq. (2) and the Supplemental Material [54]]. This is how the formation of a local moment affects the charge sector, thus representing its fingerprint. However, due to the screening effects of the bath its features get weakened, explaining the quantitative differences to the HA (e.g., the reduced size of the black square).

The most interesting situation is encountered when reducing $T$ further down to $T_{\text{low}} \gtrsim T_K$ (right panel), where the Kondo screening induces qualitative differences with respect to the HA. We observe that the low-frequency diagonal elements of $\tilde{\chi}^{\nu\nu}$ (white square) are flipped back to positive, as in the perturbative regime. This trend is driven by the low-energy correlations between electrons with antiparallel spins ($\tilde{\chi}_{\uparrow\downarrow}^{\nu\nu}$) [54]. The weakening of their negative contribution increases the physical charge susceptibility $\chi$ [see Eq. (2) and Supplemental Material [54]] and simultaneously mitigates the magnetic response. However, in the intermediate frequency regime, the diagonal elements of $\tilde{\chi}^{\nu\nu}$ are still negative, reflecting the underlying presence of a (partially screened) local moment.
The fingerprint of the Kondo regime is, thus, the onionlike frequency structure of $\tilde{\chi}_{\nu \nu}^0$, which is clearly recognizable in the rightmost central panel of Fig. 1: (i) a high-frequency perturbative asymptotic, (ii) a local moment driven structure (with suppressed diagonal) at intermediate frequencies, (iii) an inner core [with a similar sign structure as (i)] induced by the Kondo screening. A quick glance at the sign structure of $\tilde{\chi}_{\nu \nu}^0$ therefore allows for an immediate understanding of the underlying physics. This nicely illustrates the balanced competition in the charge sector between the freezing effects of the local moment and the defreezing effects of its low-energy screening, which characterizes the Kondo regime.

Note, that the onionlike structure is also found for other values of $U$, as well as in other models [54], discussed below.

How to extract the Kondo temperature.—The behavior described above is also reflected in the temperature evolution of the lowest frequency entries of $\tilde{\chi}_{\nu \nu}^0$: the diagonal $\tilde{\chi}^D = T^2 \tilde{\chi}_{T,T}^T$ and the off-diagonal $\tilde{\chi}^O = T^2 \tilde{\chi}_{-T,-T}^T$, shown in the lowest panel of Fig. 1. We can readily trace the sign changes marking the three regimes discussed above, associating the (negative) minimum of $\tilde{\chi}^D$ with the temperature at which the strongest local moment effects are observed. The screening induced enhancement of $\tilde{\chi}^D$ at lower temperatures has remarkable consequences: We find that crossing the Kondo temperature, as defined in a standard way from the behavior of the static magnetic response of the system [54] ($T_K = 1/65 \approx 0.015$ at $U = 5.75$ for the AIM), matches with high accuracy the equality of $\tilde{\chi}^D$ and $\tilde{\chi}^O$ observed at low-$T$ (see inset of Fig. 1, marked by black triangle).
Hence, \( T_\text{K} \) temperature equals the effective Kondo temperature, i.e., a cut criterion for determining all temperatures below the transition to the onset of low-energy electronic coherence: For additional Fermi-liquid behavior of the physical response can be expected [e.g., \( \rho(T) \propto T^2, c_V(T) \propto T \), etc. [11]]. This would also be consistent with the \( \tilde{\chi}^D = \tilde{\chi}^O \) condition approaching the Mott Hubbard metal-insulator transition (MIT) at \( U_{\text{MIT}}(T = 0) = U_c \) in the low-\( T \) limit (see also recent DMFT studies of the physics in the proximity of the MIT [88,89]).

The equality of the elements of the innermost \( 2 \times 2 \) submatrix of \( \tilde{\chi}^{O \nu \nu} \) represents therefore a very simple, clearcut criterion for determining \( T_\text{K} \) at the 2P level.

A nonperturbative Fermi liquid.—Beyond its physical relevance, our improved 2P understanding sheds light onto the nontrivial relation with the breakdown of perturbation theory [41]. At high \( T \), where \( \epsilon_0 = \pi T \gg V, U, t \), the \( 2 \times 2 \) submatrix encodes all relevant energy scales, the rest being nonsingular high-frequency asymptotics. In this case \( \tilde{\chi}^D = \tilde{\chi}^O \) corresponds to a singular eigenvalue of the entire \( \tilde{\chi}^{O \nu \nu} \) and hence to a divergence of the irreducible vertex function \( \Gamma^{O \nu \nu} = [\tilde{\chi}^{O \nu \nu}]^{-1} - [\tilde{\chi}_0^{O \nu \nu}]^{-1} \), specifically to the first (I) one encountered when reducing the temperature (red line in Figs. 1 and 2) [42,45,47–49]. For intermediate temperatures, the \( 2 \times 2 \) submatrix is controlled by the local moment, leading to a strongly negative \( \tilde{\chi}^D \) and negative eigenvalues of the submatrix (as in the HA case). At \( T_\text{K} \) the eigenvalue flips sign and one finds again \( \tilde{\chi}^D > \tilde{\chi}^O \) for \( T \lesssim T_\text{K} \), as in the perturbative regime (see Fig. 1, lowest panel). Here, however, because of the onionlike structure of \( \tilde{\chi}^{O \nu \nu} \), the positive definiteness (and thus the invertibility) is guaranteed only for an inner submatrix describing the Fermi liquid regime, but not for the full \( \tilde{\chi}^{O \nu \nu} \). This explains why divergences of irreducible vertex functions can occur also at low temperatures [47] even in the presence of a Fermi liquid ground state. Indeed, such vertex divergences mark the distinction between a Fermi liquid in the weak-coupling and in the strong-coupling regime.

Limitations of perturbative approaches.—The direct link between the 2P fingerprints of local moments and vertex divergences, sets precise physical limitations for perturbative methods, where—per construction—\( \Gamma \) is finite [90]. Hence, the impact of the characteristic physics emerging from the magnetic sector onto the charge channel, cannot be described by perturbative methods. We substantiate this statement by considering two advanced perturbative schemes, the functional renormalization group (fRG) [9,54] and the parquet approximation (PA) [16,54,91–101]. The results obtained for the AIM
of Fig. 1 (AIM, central): The diagonal elements are all qualitatively different from the (numerically) exact one computed by the fRG and PA (upper panels) appear with Fig. 3. Generalized charge susceptibility $\tilde{\chi}_{\nu \nu}$ computed with different approaches as a function of $T$.

with $U = 5.75$ and $T = T_{\text{int}}$ are shown in Fig. 3. $\tilde{\chi}_{\nu \nu}$ computed by the fRG and PA (upper panels) appear qualitatively different from the (numerically) exact one of Fig. 1 (AIM, central): The diagonal elements are all positive and substantially larger than the off-diagonal ones. This ensures the positive definiteness of the entire $\tilde{\chi}_{\nu \nu}$, preventing the suppression effects of the charge response, which characterize the local moment regime. This drawback qualitatively affects the physical description. In particular, the temperature dependence of the numerically exact physical charge susceptibility $\chi$ (Fig. 3, lower panel) exhibits a clear minimum for intermediate $T_{\text{high}} > T > T_K$. This emerges from the competition between the suppression induced by the local moment (see the extreme HA case) and the low-energy screening. Both features are not captured by the fRG (blue pentagons) and PA (brown squares), which display a monotonic behavior as $T$ is decreased, in the framework of a mere thermal quenching. At the same time, the perturbative approaches are able to capture the qualitative correct behavior of the magnetic response, reflecting the absence of divergences of $\Gamma$ in this sector.

Conclusions.—We have shown how fundamental physical properties of correlated systems, i.e., the local moment formation and its Kondo screening, can be directly read from the Matsubara frequency structure of the generalized charge susceptibility $\tilde{\chi}_{\nu \nu}$. In particular, the competition between localization effects at higher energies and metallic screening at lower energies is encoded in a clearly recognizable “onionlike” fingerprint of $\tilde{\chi}_{\nu \nu}$, emerging in the Kondo regime. The thorough inspection of the latter even discloses an alternative route to extract $T_K$ from the charge sector. Our improved understanding of the 2P processes sets also clear-cut limits to the physics accessible to perturbative approaches.

As a future perspective, it will be worth to overcome the on-site and/or single-orbital framework of our study. We expect that the role of the local moments will be played by short-range or Hund’s-driven magnetic fluctuations. Their nonperturbative images could reverberate, analogously as presented here, onto the charge and/or pairing response of the system. The identification of the corresponding fingerprints may open new pathways toward a microscopic understanding of unconventional superconductivity in the nonperturbative regime.

We thank M. Capone, S. Ciuchi, J. von Delft, K. Held, C. Hille, A. A. Katanin, F. Krien, F. B. Kugler, E. van Loon, C. Schattauer, and G. Sangiovanni for insightful discussions. The authors also want to thank the CCQ of the Flatiron Institute (Simons Foundation) for the great hospitality. The present work was supported by the Austrian Science Fund (FWF) through the Project No. I 2794-N35 (P. C., A. T.) and Erwin-Schrödinger Fellowship J 4266—“Superconductivity in the vicinity of Mott insulators” (SuMo, T. S.), the Deutsche Forschungsgemeinschaft (DFG) through Project No. AN 815/6-1 (S. A.), as well as the European Research Council for the European Union Seventh Framework Program (FP7/2007-2013) with ERC Grant No. 319286 (QMAC, T. S.).

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