Collinear scattering and long-lived excitations in two-dimensional Fermi gases

Serhii Kryhin, Leonid Levitov

Department of Physics, Massachusetts Institute of Technology, Cambridge, MA 02139

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Predictions of the many-body theory of Fermi-liquids are commonly viewed as trustworthy and free of significant blank spots. As a rare exception to this belief, here we argue that the seminal Landau’s $T^2$ scaling describing quasiparticle decay at low temperatures, well established in three-dimensional Fermi-liquids, undergoes a surprising change in 2D systems. We show that 2D Fermi-liquids harbor a family of abnormally long-lived excitations with lifetimes that span a wide range of time scales. It includes, in particular, excitations with lifetimes that exceed by orders of magnitude the benchmark Fermi-liquid values, with the Landau scaling replaced with $T^\alpha$, $\alpha \sim 4$. We evaluate these super-Fermi-liquid lifetimes by a direct method, establishing a link with the collinear character of quasiparticle scattering. We corroborate these results by an analytic expansion in the parameter $T/T_F \ll 1$ that employs a mapping to a 1D Schroedinger equation for a fictitious particle in a supersymmetric secular potential. Excitations with uniquely long lifetimes, resulting from quenching of the Landau damping, point to new ways of extending coherence in electron systems.

Two-dimensional (2D) metals have long been known to display quasiparticle scattering of a unique collinear character, arising due to phase space constraints for scattering at the Fermi surface[1,2]. These collinear processes are generic and largely insensitive to the specifics of two-body interactions or the details of particle dispersion. The unique quasi-one-dimensional behavior arising from these processes endows the kinetics of 2D fermions with angular memory and gives rise to peculiar ‘tomographic’ response effects[3-5]. This behavior calls for a comparison with one-dimensional (1D) systems, where collinear scattering renders quasiparticles short-lived, destroying the Fermi-liquid state and replacing it with the Tomonaga-Luttinger state[6,7]. Below we argue that collinear processes in 2D metals take on a role which is a complete opposite of that in 1D liquids. These processes give a giant boost to quasiparticle lifetimes and can be said to produce a “super-Fermi-liquid” that harbors a unique family of excitations with exceptionally long lifetimes, exceeding by orders of magnitude those familiar from Fermi-liquid theory. The extinction of the Landau $T^2$ damping for these excitations points to new interesting ways for extending coherence in electron systems.

The occurrence of new time scales becomes particularly transparent in a 2D metal with an isotropic particle dispersion and circular Fermi surface, since in this case different excitations are associated with different angular harmonics of Fermi surface modulations evolving in space and time. In that, the abnormally long-lived excitations are identified with the odd-$m$ angular harmonics, whereas the even-$m$ harmonics feature conventional Fermi-liquid lifetimes. As illustrated in Fig.4 at low temperatures $T \ll T_F$ these long lifetimes greatly exceed those in Fermi-liquids and show strong departure from conventional scaling. The decay rates in Fig.4 are obtained by a direct calculation that treats quasiparticle scattering exactly, using a method that does not rely on the small parameter $T/T_F \ll 1$. The odd-$m$ decay rates display scaling $\gamma \sim T^\alpha$ with super-Fermi-liquid exponents $\alpha > 2$. In our analysis we find $\alpha$ values close to 4, i.e., the odd-$m$ rates are suppressed strongly compared to the even-$m$ rates, $\gamma_{odd}/\gamma_{even} \sim (T/T_F)^2$.

These findings may seem to contradict the well-known results for excitation lifetimes in 2D Fermi gases found from Green’s function selfenergy calculations, which predict that collinear scattering shortens quasiparticle lifetimes[8-14]. Namely, decay rates predicted in this way are faster by a log factor $\log(T_F/T)$ than the conventional $T^2$ rates. The selfenergy approach is therefore conspicuously unaware of the existence of the long-lived odd-$m$ excitations. This may seem surprising also because it is usually taken for granted that there is a single timescale that characterizes decay for all low-energy excitations. However, as made clear by Fig.4, this is very much untrue...
in 2D, since the odd- and even-m modes have drastically different lifetimes that show different scaling vs. \( T \). The conventional selfenergy approach is not well suited for such a situation, since selfenergy is the quantity which is most sensitive to the fastest decay pathways. It is probably for this reason that the long-lived excitations, the main finding of the present work, have been missed in the literature despite the 60 years of intense interest in Fermi liquids.

We also emphasize that, similar to the self-energy analysis\[^8\]–\[^14\], the occurrence of long-lived excitations is a robust property that persists for non-circular Fermi surfaces, so long as the distortion away from the circle is not big. The reason is that the inversion symmetry, whenever present, separates modulations of the Fermi surface into modes of an even and odd parity. The difference in lifetimes for these mode types is identical to that found here for circular Fermi surfaces.

It is interesting to note that in some electron systems collinear dynamics can speed up quasiparticle decay rather than slow it down. Speed-up arises because collinear scattering, by allowing particles to travel side by side for a longer time and thereby interact more strongly, enhances the effective interactions and shortsens lifetimes. This physics is well documented, e.g., for Dirac bands, where collinear dynamics arising from linear band dispersion impacts carrier lifetimes and dynamics\[^\ref{15}\]–\[^22\]. In our problem, a different behavior originates from the delicate interplay of collinear scattering and phase space constraints. These effects dominate at a 2D Fermi surface but are of little importance for highly excited states in Dirac bands.

One more reason for why the long-lived modes have been missed in the literature undoubtedly lies in the difficulty of a direct calculation. To gain insight and understand better the origin of this difficulty we consider excitations at the Fermi surface in the framework of the Fermi-liquid kinetic equation. An ingenious approach developed in Refs.\[^23\]–\[^26\] allows to tackle the kinetic equation, linearized near thermal equilibrium at \( T < T_F \), bringing it to the form of a time-dependent Schroedinger equation with a reflectionless seanch potential. This equation, being exactly soluble, successfully predicts a \( T^2 \) scaling for excitation decay in 3D. However, it fails when applied in 2D, yielding unphysical vanishing rates.

It is instructive to inspect more closely the exactly-solvable kinetic equation approach\[^23\]–\[^26\] and try to understand why it fails in 2D case. The starting point is the Fermi-liquid transport equation

\[
\frac{df_i}{dt} + [f, H] = \sum_{21'2'} (w_{1'2'\rightarrow 12} - w_{12\rightarrow 1'2'}),
\]

where \( f(p, r, t) \) is fermion distribution, \([f, H]\) denotes the Poisson bracket \( \nabla_r f \nabla_p e - \nabla_r e \nabla_p f \). The right-hand side is the rate of change of the occupancy of a state \( p_1 \), given as a sum of the gain and loss contributions resulting from the two-body scattering processes \( 12 \rightarrow 1'2' \) and \( 1'2' \rightarrow 12 \). Fermi’s golden rule yields

\[
w_{1'2' \rightarrow 12} = \frac{2\pi}{\hbar} |V_{12'12}|^2 \delta(p - p_1)(1 - f_1)(1 - f_2) f_{1'2'},
\]

where the delta functions \( \delta = \delta(e_1 + e_2 - e_{1'2'}) \), \( \delta_p = \delta(p_1 + p_2 - p_{1'2'}) \) account for the energy and momentum conservation. The gain and loss contributions are related by the reciprocity symmetry \( 12 \leftrightarrow 1'2' \). Here \( V_{12'12} \) is the two-body interaction, properly antisymmetrized to account for Fermi statistics. Interaction \( V_{12'12} \) depends on momentum transfer \( k \) on the \( k \sim k_F \) scale; this \( k \) dependence is essential and will be ignored. In what follows we consider a spatially uniform problem setting \([f, H] = 0\). The sum over momenta \( 2', 1' \) represents a six-dimensional integral over \( p_2, p_1 ', p_2' \), which is discussed below.

For the states weakly perturbed away from equilibrium, \[^2\] can be linearized using the standard ansatz \( f(p) = f_0(p) - \frac{2\eta}{\hbar} \eta(p) \). After standard algebra, this yields a linear integro-differential equation

\[
\frac{df_0}{dt} + [f_0, H] = I_{ee} \eta,
\]

\[
I_{ee} \eta = \sum_{21'2'} \frac{2\pi}{\hbar} |V|^2 F_{12'12} \delta_p (\eta_{1'} + \eta_{2'} - \eta_1 - \eta_2).
\]

Here \( \sum_{21'2'} \) and \( |V|^2 \) denote the six-dimensional integral \( \int d^6p d^6p_1 d^6p_2 \) and the interaction matrix element \( |V_{12'12}|^2 \), the quantity \( F_{12'12} \) is a product of the equilibrium Fermi functions \( f^0_{12} - f^0_{12'} \). Different excitations are described by eigenfunctions of the collision operator \( I_{ee} \), with the eigenvalues giving the decay rates equal to inverse lifetimes. Because of the cylindrical symmetry of the problem, the eigenfunctions are products of angular harmonics on the Fermi surface and functions of the radial energy variables \( x_i = \beta(\epsilon - \mu) \):\(^3\)

\[
\eta(p, t) = e^{-\gamma_m t} e^{i\epsilon t} \chi_m(x),
\]

where \( \gamma_m \) and \( \chi_m(x) \) are solutions of a spectral problem

\[
-\gamma_m f_0(1 - f_0) \chi_m(x) = I_{ee} \chi_m(x).
\]

In general, the six-dimensional integral operator \( I_{ee} \) has a complicated structure which is difficult to analyze. However, at \( T \ll T_F \) the part of phase space in which transitions \( 12 \leftrightarrow 1'2' \) are not restricted by fermion exclusion is a thin annulus of radius \( p_F \) and a small thickness \( \delta p \approx T_F / v \ll p_F \). One can therefore factorize the six-dimensional integration over \( p_2, p_1 ', p_2' \) in \( I_{ee} \) into a three-dimensional energy integral and a three-dimensional angular integral, and integrate over angles to obtain a close-form equation for the radial dependence \( \chi(x) \). This is done by noting that the delta functions \( \delta_p \) together with the conditions \( |p_1| \approx |p_2| \approx |p_1'| \approx |p_2'| \approx p_F \) imply that the states \( 1, 2, 1' \) and \( 2' \) form two anti-collinear pairs

\[
p_1 + p_2 \approx 0, \quad p_1 ' + p_2 ' \approx 0
\]
The azimuthal angles therefore obey \( \theta_1 \approx \theta_2 + \pi, \theta_1' \approx \theta_2' + \pi \). In a thin-shell approximation \( \delta p \ll p_F \), this gives two delta functions \( \delta(\theta_1 - \theta_2 - \pi), \delta(\theta_1' - \theta_2' - \pi) \) that cancel two out of three angle integrals in \( I_{\text{ex}} \), allowing to rewrite the quantity \( \eta_1' + \eta_2' - \eta_1 - \eta_2 \) as
\[
e^{im\theta_1'}(\chi(x_1')) + (-)^m\chi(x_2')) - e^{im\theta_1}(\chi(x_1) + (-)^m\chi(x_2)).
\]
(6)

Subsequent steps differ for even and odd \( m \), because the contributions of \( \chi(x_1') \) and \( \chi(x_2') \) to \( I_{\text{ex}} \) cancel out for odd \( m \) and double for even \( m \), since \( F \) is symmetric in \( x_1' \) and \( x_2' \). Focusing on odd \( m \) and carrying out integration over the angle between \( \mathbf{p}_1 \) and \( \mathbf{p}_1' \) yields
\[
\tilde{F} \frac{d\chi(x_1)}{dt} = T^2 \int dx_2 dx_1 dx_2' F \delta_x[\chi(x_1) - \chi(x_2')],
\]
(7)
where \( \tilde{F} = f_0(1 - f_0) \) and \( \delta_x = \delta(x_1 + x_2 - x_1' - x_2') \). Here \( T^2 \) originates from nondimensionalizing the energy variables \( x_i \) in the integral and the delta function, the dimensionless factor \( g \) is a result of angular integration, the quantity \( F \) is defined above. Integration over energy variables \( x_2, x_1', x_2' \) extends throughout \(-\infty < x_i < \infty \), as appropriate for \( T < T_F \).

This equation, after introducing new distribution as \( \chi(x) = 2\cosh \frac{x}{2} \zeta(x) \) and carrying out Fourier transform in the energy variable, \( \zeta(x) = \int dk e^{ikx} \psi(k) \), can be rewritten as a time-dependent Schrödinger equation for a particle moving in a one-dimensional scissant potential
\[
\partial_t \psi(k) = g T^2 \left[ \frac{\pi^2}{\cos \pi k} - \psi(k) - \frac{1}{2} \psi''(k) \right],
\]
(8)
(see Supplement). Unlike the 3D case, where after a similar transformation the \( T^2 \) scaling translates into a \( T^2 \) dependence of the decay rates, here the operator in (8) has a zero mode, \( \psi_0(k) = \frac{1}{\cosh(\pi k)} \). Being a zero mode, this mode does not relax. The associated \( \chi_0(x) \) can be found from the identity \( \int d\xi e^{i\pi \xi} = \frac{\pi}{\tan \pi k} \), giving \( \chi_0(x) = 1 \). Returning to the energy variable, this yields the Fermi-surface-displacement mode \( \delta f(x) = df_0/df = f_0(1 - f_0) \), identical for all odd \( m \).

For even \( m \), analysis proceeds in a similar manner, however it yields a normal \( T^2 \) scaling of the decay rates. This is so because for even \( m \) different terms in (6) are of equal signs and do not cancel out. As a result, the lifetimes for the even-\( m \) and odd-\( m \) harmonics are quite different. To capture this difference without running into the unphysical infinite lifetimes, the zero-thickness approximation for the active shell at the Fermi surface must be relaxed. This is difficult to do in the framework described above and one must seek alternative ways.

Here we proceed in two steps, first using the kinematic constraints to reduce the six-dimensional integral in (5) to a three-dimensional integral, and then using a suitable basis of functions to reduce the three-dimensional integrals to one-dimensional integrals. After that, the operator \( I_{\text{ex}} \) can be projected on a subspace that represents adequately the states on the active shell and diagonalized numerically.

Integration over \( p_2 \) can be eliminated by a momentum-conservation delta-function, giving
\[
I[\eta] = -\frac{2\pi}{\hbar} |V|^2 \int dp_1' dp_2' \delta_{\epsilon_1'} \sum_{\alpha} \eta_\alpha,
\]
where \( I[\eta] \) is now a function of the other momenta, \( p_2 = p_1' + p_2' - p_1 \). As above, \( \delta_{\epsilon_1'} \) denotes \( \delta(\epsilon_1' - \epsilon_2 - \epsilon_1 - \epsilon_2) \) and \( \sum_{\alpha} \eta_\alpha \) stands for \( \eta_1' + \eta_2' - \eta_1 - \eta_2 \). Next, we eliminate the radial integration over \( |p_2'| \) by canceling it with the energy delta-function. The expression for the collision integral then takes the form
\[
I[\eta] = -A \int dp_2 \frac{d\theta_n}{(2\pi)^2} F_{121'2'} \delta_{\epsilon_2'} \sum_{\alpha} \eta_\alpha, 
\]
(10)
where we introduced a temperature-independent constant \( A = \pi m |V|^2 / \hbar^3 \).

Due to momentum and energy conservation, in (10) the momenta \( p_{1'} \) and \( p_{2'} \) satisfy the constraints
\[
p_{1'} = p_+ + |p_-| n, \quad p_{2'} = p_+ - |p_-| n,
\]
(11)
where \( p_{\pm} = p_{\pm} - p_0 \) and we introduced a unit vector \( n = (\cos \theta_n, \sin \theta_n) \) that parameterizes the outgoing momenta in the collision process, wherein the incoming momenta \( p_1 \) and \( p_2 \) are taken to be fixed, see schematic in Fig. 2.

Next, we choose a basis of functions to represent the states \( \eta(p_{1'}) \) and define a matrix representation for the linear operator \( I[\eta_{p_1}] \). Different choices of basis functions have different computational limitations. Here we employ, as a basis, the \( \delta \)-functions
\[
\eta_k(p) = \delta^{(2)}(p - k)
\]
(12)
cumbersome. In the term with the delta functions, (12), when substituted in the collision operator, (10), the collision operator can be simplified, as above, by canceling two out of three integrations in \( \int d\rho_2 d\theta_2 d\theta_n \ldots \) yielding an expression that involves just one integral.

An added benefit of working in the delta-function basis is that it allows to analytically simplify the expression for the collision operator, (10). The collision operator can be written as a sum of four contributions, one for each \( \eta_n \). This yields an expression \( I[\eta] = I_1[\eta] + I_2[\eta] - I_3[\eta] - I_4[\eta] \) with the individual terms given below:

\[
I_1[\eta_k] = -A \eta_k(p_1) \int \frac{d^2 p_2 d\theta_n}{(2\pi)^4} F_{121'2'},
\]

\[
I_2[\eta_k] = -A \int \frac{d\theta_n}{(2\pi)^4} F_{121'2'},
\]

where \( I_1 \) and \( I_2 \) represent the contributions of \( \eta_1 \) and \( \eta_2 \) in \( [10] \) and \( F \), as above, denotes \( F = f_1^0 f_2^0 (1 - f_1^0) (1 - f_2^0) \). In \( [14] \) we eliminated two integrals by integrating over a delta-function. Integrals \( I_3 \) and \( I_4 \), which correspond to \( \eta_1 \) and \( \eta_2 \) respectively, can be written in a similar way:

\[
I_3[\eta_k] = -A \int \frac{d^2 p_2 d\theta_n}{(2\pi)^4} F_{121'2'} \eta_k(p_{1'v}),
\]

\[
I_4[\eta_k] = -A \int \frac{d^2 p_2 d\theta_n}{(2\pi)^4} F_{121'2'} \eta_k(p_{2'})
\]

In this case, eliminating integration over \( p_2 \) by canceling it with the delta functions \( \eta_k(p_{1'v}), \eta_k(p_{2'}) \) is a little more cumbersome. In the term \( I_1 \) the \( \delta \)-function constraint is

\[
p_{2'}(p_1, p_2, \theta_n) = k,
\]

where the expression for \( p_{2'} \) is given in \( [11] \). This equation should be solved for \( p_{2'}(p_1, k, \theta_n) \), which is a zero of the \( \delta \)-function’s argument. To perform integration over \( p_2 \) in \( [16] \) we use the value \( p_{1'}(p_1, k, \theta_n) \) and evaluate the Jacobian at the zero of the delta function. Conveniently, Eq. \( [17] \) can be solved in a closed form, after which the first relation in \( [11] \) yields

\[
p_{2'}(p_1, k, \theta_n) = 2k - p_1 - \frac{(k - p_1)^2 n}{(k - p_1) \cdot n}
\]

\[
p_{1'}(p_1, k, \theta_n) = k - \frac{(k - p_1)^2 n}{(k - p_1) \cdot n}
\]

The Jacobian of a \( \delta \)-function in \( I_4 \) is

\[
J = \frac{(\partial p_{2'v}, \partial p_{2'})}{(\partial p_{1'v}, \partial p_{1'})} = \frac{1}{2} \frac{(k - p_1) \cdot n}{(k - p_1)^2}
\]

(found by linearizing the second relation in \( [11] \)). After handling the contribution \( I_3 \) in a similar manner, the sum of \( I_3 \) and \( I_4 \) can be simplified to read

\[
(I_3 + I_4)[\eta_k] = -A \int \frac{d\theta_n}{(2\pi)^4} J^{-1} F_{121'2'},
\]

where \( p_2 \) and \( p_{1'} \) are given by Eqs. \( [18] \) and \( [19] \), and the Jacobian \( J \) is given by \( [20] \).

Importantly, evaluating \( I_2, I_3 \) and \( I_4 \) on a delta function state, Eq. \( [11] \) yields smooth functions of \( p \) given by simple 1D integrals. The \( I_1 \) contribution, to the contrary, yields a delta function identical to the one in \( [12] \), with a prefactor that is given by a 3D integral. This contribution describes particle loss from the initial state \( p_1 \), the contributions \( I_2, I_3 \) and \( I_4 \) describe gain.

One peculiar aspect of working with delta functions is a Jacobian that has a non-analytic structure, \( [20] \). We note...
that, while the Jacobian $J$ is zero when condition $(k - p_1) \cdot n = 0$ is satisfied, this does not mean that the whole expression inside the integral is divergent. The behavior of the integral around the divergent points of a phase space can be understood by introducing new variables $\Delta p$ and $\phi$, such that $\Delta p = |k - p_1|$ and $\cos \phi = (k - p_1) \cdot n/|\Delta p|$. In these variables the Jacobian $J$ can be written as

$$J = \cos^2 \phi,$$

(22)

an expression that remains finite and non-zero so long as $\cos \phi \neq 0$. Therefore, a divergence in the integral in (21) might occur only when the quantity $\cos \phi$ vanishes. On the other hand, expressions for $p_2$ and $p_1'$ in Eqs. (18) and (19) have a term $(p_1 - k) \cdot n$ in their denominators, which is proportional to $\cos \phi$. Therefore at $\cos \phi = 0$ the absolute values of $p_2$ and $p_1'$ diverge so that $|p_2| \to +\infty$ and $|p_1'| \to +\infty$. This divergence leads to an exponential decrease of the $f(p_2)$ term, which cancels the divergence of $J^{-1}$. Therefore, the expression inside the integral has only an isolated discontinuity point rather than a pole and therefore the integral has a finite value. Even though the integral is always integrable, the singularity near $\mathbf{p} = \mathbf{k}$ point makes the numerical computation problematic. We study the impact of the numerical error in forward scattering in the supplement and find out that it does not affect the qualitative behavior.

The representation of the collision operator introduced above can be used to project it on a subspace spanned by a set of basis functions chosen to provide a sufficiently good sampling of the active region in momentum space (the blurred annulus pictured in Fig.2). This yields a finite-size matrix that can be diagonalized to find the excitation eigenmodes and their eigenvalues, giving the decay rates. We have found that, although this direct approach works, it is more convenient to use a somewhat different approach to the problem, which employs the angular distribution of quasiparticle scattering in the active region near the Fermi surface. The angular distribution, besides being directly linked to the individual excitation modes and their lifetimes, as discussed below, also allows to establish an interesting connection to the excitations with di

$$\gamma_m \text{ are related to the Fourier coefficients of the angular distribution,}$$

(24)

$$\sigma(\theta) = \sum_m e^{im(\theta - \theta_0)}(\gamma_m - \gamma_0),$$

where the term $-\gamma_0$ describes particle loss from the injected beam. We use the basis functions introduced above to compute $\sigma(\theta)$ and then use the relation in (21) to obtain lifetimes of different modes.

The angular dependence, shown in Fig.3, features sharp peaks centered at $\theta = 0$ and $\pi$ with the angular widths that scale as $T \ll T_F$, describing forward scattering and backscattering, respectively. Interestingly, the backscattering peak is of a negative sign, representing backreflected holes. At $T \approx T_F$ the values $\sigma(\theta)$ at generic $\theta$ are found to scale as $T^2/T_F$, as expected from Fermi-liquid theory. This behavior is detailed in Fig.3 insets. Despite the overall $T^2$ scaling, the decay rates $\gamma_m$ for the odd-$m$ modes, found from the relation in (24), show significant departure from the $T^2$ scaling. The decay rates for harmonics $e^{im\theta}$ with even and odd $m$, shown in Fig.1, are similar at $T \sim T_F$ but show a very different behavior at $T < T_F$. This difference between even-$m$ and odd-$m$ rates originates from the collinear character of scattering, manifest in the strong peaks in $\sigma(\theta)$ in the forward and backward directions. The nearly equal areas of these peaks and the negative sign of the backscattering peak suppress the odd-$m$ Fourier harmonics of $\sigma(\theta)$, giving small decay rates for these harmonics. The temperature dependence for the even-$m$ harmonics agrees well with the $T^2$ law. The odd-$m$ harmonics, to the contrary, have decay rates decreasing at low $T$ much faster than $T^2$. For these harmonics, the observed scaling is $\gamma_m \sim T^n$ with $n$ slightly below 4, which can be described as a “super-Fermi-liquid” suppression of the decay rates for odd-$m$ harmonics.

Lastly, it is interesting to mention that collinear scattering, manifest in the sharp peaks in the angle-resolved cross-section $\sigma(\theta)$ at $\theta = 0$ and $\pi$, is directly responsible for the log enhancement of quasiparticle decay rates predicted from the self-energy analysis. Indeed the angle dependence near $\theta = 0$ and $\pi$ is of the form $\sigma(\theta) \sim T^2/|\theta|$ and $T^2/|\theta - \pi|$, with the $1/|\theta|$ singularity rounded on the scale $\delta \theta \sim T/T_F$, as illustrated in Fig.3 (see above). Integrating the angle-resolved crosssection over $\theta$ yields a $\log(T_F/T)T^2$ total scattering crosssection.

This illustrates that the abnormally long lived excitations with the decay rates that scale as $T^4$ rather than $T^2$, described in this work, and the seminal $\log(T_F/T)T^2$
decay rates, originate from the same phase-space constraints. Restricted phase space renders quasiparticle scattering a highly collinear process even when the microscopic interactions have a weak angular dependence. The unusual kinetics originating in this regime, is relevant for a variety of 2D systems, in particular those where small carrier density and small kinetic energy make electron-electron collisions a dominant scattering mechanism that overwhelms other carrier relaxation pathways.

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Supporting Material for “Collinear scattering and long-lived excitations at a 2D+++ Fermi surface”

Transforming the collision integral to the partial differential equation

In this section we show the transformation of the collision integral into a second order partial differential equation, which can be carried out after the integration over angles is separated from the integration over energies.

As a first step, we reverse signs of the $1'$ and $2'$ variables: $x_{1'} \to -x_{1'}$, $x_{2'} \to -x_{2'}$. This transforms the integral equation Eq. 7 to

$$
\tilde{F} \frac{d\chi}{dt} = gT^2 \int dx_2 dx_{1'} dx_{2'} F_{121'} \delta^+_x (\chi(x) - \chi(x_2))
$$

$$
F_{121'} = f_0(x_1) f_0(x_2) f_0(x_{1'}) f_0(x_{2'})
$$

(25)

where $\delta^+_x = \delta(x_1 + x_2 + x_{1'} + x_{2'})$. Next we use the identities

$$
\int dx_2 dx_{1'} dx_{2'} f_0(x_2) f_0(x_{1'}) f_0(x_{2'}) \delta^+_x = \frac{1}{2} \frac{x_1^2 + \pi^2}{1 + e^{-x_1}},
$$

(26)

$$
\int dx_1 dx_{2'} f_0(x_{1'}) f_0(x_{2'}) \delta^+_x = -\frac{x_1 + x_2}{1 - e^{-x_1 - x_2}}.
$$

(27)

to carry out integration over $x_2, x_{1'}, x_{2'}$ in the first term and over $x_{1'}, x_{2'}$ in the second term. This brings the integral equation to the form

$$
\tilde{F} \frac{d\chi}{dt} = gT^2 \left[ f_0(x_1) \frac{1}{2} \frac{x_1^2 + \pi^2}{1 + e^{-x_1}} \chi(x) + \int dx_2 f_0(x_2) \frac{x_1 + x_2}{1 - e^{-x_1 - x_2}} \chi(x_2) \right]
$$

(28)

(29)

This equation can be simplified using the substitution

$$
\chi(x) = 2 \cosh \left( \frac{x}{2} \right) \zeta(x) = \left( e^{x/2} + e^{-x/2} \right) \zeta(x),
$$

(30)

which gives an equation

$$
\frac{d\zeta(x_1)}{dt} = gT^2 \left[ \frac{x_1^2 + \pi^2}{2} \zeta(x_1) + \int dx_2 \frac{x_1 + x_2}{2 \sinh \frac{x_1 + x_2}{2}} \zeta(x_2) \right]
$$

Next, we reverse the sign of $x_2$, which brings the integral operator to the form of a convolution, separately for the even and odd functions $\zeta(x_2)$. For an even function $\zeta(-x_2) = \zeta(x_2)$ we have

$$
\int dx_2 \frac{x_1 - x_2}{2 \sinh \frac{x_1 - x_2}{2}} \zeta(x_2).
$$

After Fourier transform $\zeta(x) = \int dk e^{ikx} \psi(k)$ this gives the time-dependent Schrödinger equation with a secanth potential $\frac{\sinh^2 x}{\cosh^2 x}$ displayed in the main text,[31]. Physical solutions correspond to the eigenfunctions that are even in $x$. The zero mode $\psi_0(k) = \frac{1}{\cosh(\pi k)}$, being an even function of $k$, is part of this family. Going back to the $x$ variable and undoing the substitutions gives the zero mode $\chi_0(x) = 1$ for all odd $m$, and $\delta f_0(x) = f_0(x)(1 - f_0(x))$ as discussed in the main text.

Analogously, for odd functions $\zeta(-x_2) = -\zeta(x_2)$ upon changing $x_2$ to $-x_2$ a minus sign appears in front of the integral operator:

$$
-\int dx_2 \frac{x_1 - x_2}{2 \sinh \frac{x_1 - x_2}{2}} \zeta(x_2).
$$
Carrying out Fourier transform $\zeta(x) = \int dk e^{ikx} \psi(k)$ give a time-dependent Schroedinger equation for a particle moving in a secant potential of an opposite sign

$$\partial_t \psi(k) = gT^2 \left( \left( \frac{\pi^2}{2} + \frac{\pi^2}{\cosh^2 \frac{\pi}{k}} \right) \psi(k) - \frac{1}{2} \psi''(k) \right)$$ (31)

In this case, physical solutions correspond to the eigenfunctions that are odd in $k$. For a repulsive secant potential these functions are in the continuum spectrum and asymptotically have the form of plane waves. This indicates that the behavior of the eigenfunctions that are odd in $x$ differs from that of the even eigenfunctions discussed above. This poses interesting questions for future work.

The delta-function basis and an optimized integration mesh

Here we discuss the basis of functions in the momentum space used to represent particle momentum distributions perturbed by collisions. We describe the reasoning behind our basis choice and the integration mesh used in this study. In this work we opted for a basis comprised of suitably normalized delta functions. This choice is quite different from the more conventional approaches relying on systems of orthogonal polynomials (e.g. see [4]). The delta functions, being singular functions, may not appear to be a natural choice of a basis. However, in a problem like ours, the delta functions have distinct advantages, since, after being plugged in Eq.(10) of the main text they considerably reduce the number of required integrations.

As we observed, the nature of the $\delta$-functions allows to eliminate two out of three integrations in three out of four terms in plugged in Eq. (eqnumber) of the main text. In addition, projection operation is effectively reduced to computing a value of the function of interest in the corresponding point, unlike in continuous functions bases, where we need to compute one more integral to perform the projection on the basis function itself. Additional complication comes from nature of the function inside the integral. At low temperatures it resembles several peaks in its variable space. This makes the Monte Carlo approach to the integration very hard to apply and pushes us to a mesh-based definite integration methods. Let us assume we perform the integration with $N$ mesh points in a mesh in angular variables. The computation time of the matrix element in continuous basis takes $O(M^2N^3)$ time. A non-diagonal element of the matrix, which is formed by only $I_2$, $I_3$, and $I_4$, in some $\delta$-function basis takes just $O(N)$ time. A diagonal element of a $\delta$-function basis, where $I_1$ also has an impact, takes $O(MN^2)$. To solve a linear problem in a $\delta$-function basis, we do not need to compute the matrix elements for each mesh point. As we show below, rotational symmetry of the initial expression allows us to perform the integration on $M^2N$ mesh points instead of $M^2N^2$ for any non-diagonal matrix element and $M$ points instead of $MN$ points for diagonal elements. As takeaway, rotational symmetry of the initial expression allows to circumvent one of the integrations over $\theta$. Therefore, the total computation complexity with the $\delta$-function mesh is $O(M^2N^2)$ instead of $O(M^2N^3)$ for continuous bases.

With a very specific choice of a smooth basis for this particular problem it is possible to construct a numerical solution of a the same computational complexity. In particular, one needs to be rigorous in choosing the basis in the way that the choice would respect both rotational symmetry and properties of the integral. One of the possible ways to construct such basis is to use the form

$$f_{nm}(p) = P_n(p)e^{im\theta}\hat{F}(p),$$ (32)

where $p$ and $\theta$ are polar coordinates of $p$, $\hat{F}(p) = f(1 - f)$, and $P_n(p)$ is a polynomial of power $n$ which is chosen to make $f_{nm}(p)$ to be orthogonal to $f_{n'm'}(p)$ when $n \neq n'$ with respect to the integral inner product. This basis is one of the optimal bases, since it is both rotationally invariant and spans the region around the Fermi surface, but we stick to less complex basis of $\delta$-functions.

Since computation of the collision integral on the delta-functions is much faster, we will use a sampling of delta-functions as a subspace basis for computation. We define a set of basis vectors as Kronecker $\delta$-functions

$$|p_i\rangle = \hat{\delta}^{(2)}(p - p_i)\sqrt{\Delta V_i},$$ (33)

where $\Delta V_i = p_i^2\Delta p_i\Delta\theta_i\Delta p_i$ and $\Delta\theta_i$ are sizes of the part of momentum space that corresponds to $i$'th point in polar coordinates. By $\hat{\delta}^{(2)}(p - p_i)$ in (33) we mean a a function defined on the mesh and that is equal to 0 when $p \neq p_i$, and is equal to $1/\Delta V_i$ when $p = p_i$. In the limit of dense mesh function $\hat{\delta}^{(2)}(p - p_i)$ behaves like a Dirac $\delta$ function. The square root of phase space element is added to preserve the normalization of the basis to be $\langle p_i|p_j\rangle = \delta_{ij}$ with respect to the inner product in the form of an integral over $p$. 
We use this basis to represent the operator $I$ as a matrix:
\[
\langle p_i | I | p_j \rangle \equiv I_{ij} = I(\delta^{(2)}(p_i - p_j)) \sqrt{\Delta V_i \Delta V_j}.
\]
(34)

By this construction, the expression yields a symmetric matrix. As such it is suitable for computing the angular distribution for two-body scattering, for which the matrix should be applied to a state that represents the incoming state. It should be noted, however, that the eigenvectors and eigenvalues describing different excitations and their lifetimes are not those of the matrix $I$. Rather, they should be obtained from a generalized eigenvalue problem $\gamma \tilde{F}(p)\psi = I\psi$ with $\tilde{F}(p) = f_0(1 - f_0)$.

On the side, the lowest eigenvalues for each $m$ can be determined more easily from the angular distribution, as discussed in the main text. This approach was used to obtain the eigenvalues shown in Fig. 5 and Fig. 3 of the main text. We verified that the direct solution of the generalized eigenvalue problem gives the same eigenvalues, albeit with a lower accuracy.

Next, we discuss another crucial aspect of our analysis — sampling of the relevant part of the momentum space. This achieved by constructing a mesh of points on which the delta-function states given in (33) are centered. The mesh must have a higher density near the Fermi surface and for near-collinear momenta, and also respect the cylindrical symmetry of the problem. There are several ways through which these requirements can be satisfied. Below we described the approach that proved particularly useful.

To preserve the rotational invariance of the collision operator, we take the the mesh points on a set of concentric circles centered at $\mathbf{p} = 0$, as illustrated in Fig. 4. The radial momentum components form an equally spaced set of $M$ points in an interval $p_{\text{min}}(T) < p < p_{\text{max}}(T)$ centered at $p = p_F$. To optimize coverage of the phase space within thermally broadened Fermi surface we used temperature dependence of $p_{\text{min}}(T)$ and $p_{\text{max}}(T)$ was optimized defined by $\tilde{F}(p_{\text{max}}, T) = \tilde{F}(p_{\text{min}}, T) = \alpha$, with $p_{\text{min}} < p_{\text{max}}$ and $\alpha$ a small parameter of choice. In this study we used several values of $\alpha$ and $M$ and came to conclusion that the best choice that allows to achieve reasonable precision is $\alpha = 10^{-3}$ and $M = 40$. In cases where there was no lower-limit solution for $p_{\text{min}} > 0$, the value $p_{\text{min}}$ was set to 0. The choice of boundaries on the absolute values of momentum in the mesh allows us to focus on the physically interesting region of the phase space near a Fermi surface where $\tilde{F}(p, T)$ is not exponentially small. For the temperatures $T \sim \epsilon_F$, the sampled region was a disc of the radius $\sim \epsilon_F$. For the temperatures $T \ll \epsilon_F$, the sampled region was an annulus of radius $\epsilon_F$ and thickness of $\sim T$.

We choose a specific mesh point distribution to resemble the properties of the integral as a function of the angle between momenta $p_i$ and $p_j$. To perform the collision operator analysis as a function of the angle, we need to be able to integrate over an absolute value of momentum (i.e. sum over points with the same angular coordinate and different radial coordinates). Because of this, we choose the same angular distribution of points for each circle of constant momentum absolute value. Assuming the $\delta$-function source, we find that at low temperatures most of the scattering is either near-forward scattering or near-back scattered, and the width of the forward and backward peaks scales $\sim T$ at small temperatures. To describe this highly anisotropic scattering it is beneficial to define mesh that has a higher density for the angles in the near-forward and near-backward directions. To construct a mesh with such properties we choose a uniform mesh in angle to account for the general properties of the angular distribution. To that end, we use a combination of a uniform mesh for the angles away from the colinear and anticolinear directions $\theta = 0$ and $\pi$ and a denser mesh concentrated in the regions near $\theta \approx 0$ and $\theta \approx \pi$. The width of these two regions is taken to be a function of temperature proportional to $T$, which accounts for the forward and backward scattering distribution becoming sharper as $T$ decreases, as illustrated in Fig. 3 of the main text. The dense forward/backward mesh is taken to be uniform, comprised of $N$ points. The not-so-dense mesh for non-collinear angles is also taken to be uniform, comprised of $N'$ points. This is illustrated in Fig. 4 where the dense and less dense meshes are shown in different colors. In our simulation we used $N = N' = 200$.

For a 2D mesh in momentum space we use a direct product of the radial and angular meshes defined as described above. We denote the mesh points as $|p_{mn}\rangle \equiv |p_m, \theta_n\rangle$, where $p_{mn} = (p_m \cos \theta_n, p_m \sin \theta_n)$ and $1 < m < M$ and $1 < n < N + N'$.

**Matrix representation of the linearized collision operator**

Here we describe in details the method to obtain the angular distribution from the operator projected on the functional basis and show the correspondence of these operations to the operations with original collision integral in the function space.

We choose the source in the form of delta function in an angular space to describe the electron injection along $\theta = 0$. The radial distribution for injected electrons is chosen to be proportional to $-\partial f_0/\partial \epsilon$. This corresponds to $\eta_0(p_i, \theta_j) \equiv \eta_0(\theta_j) = \delta(\theta - \theta_j)$ when $\theta_j = 0$. To focus on the angular part of the operator, we contract the matrix
with a column corresponding to $\eta_0$; this is equivalent to integrating original expression over the radial coordinates of momenta:

$$\langle \theta_i | I | \theta_j \rangle = \sum_{p,p',\theta,\theta'} \langle \eta_0(\theta_i) | p, \theta \rangle \langle p, \theta | I | p', \theta' \rangle \times \langle p', \theta' | \eta_0(\theta_j) \rangle$$  \hfill (35)

Here the summation over $p$ is a summation over all values of $p_m$ where $1 \leq m \leq M$, and summation over $\theta$ is a summation over all values of $\theta_n$ where $1 \leq n \leq N + N'$. The summation over $p$ and $\theta$ in (35) corresponds to the integration over momentum space in the following way:

$$\sum_{p,\theta} \langle \eta_0(\theta_i) | p, \theta \rangle \langle p, \theta | I | p_m, \theta_n \rangle \leftrightarrow \int_0^{+\infty} dp \int_0^{2\pi} d\theta \eta_0(\theta_i) I[\delta^{(2)}(p - p_{mn})],$$ \hfill (36)

Analogous relations can be established for summations over $p'$ and $\theta'$.

Note that $\langle \theta_i | I | \theta_j \rangle$ depends only on the angles and does not depend on the absolute values of two momenta anymore. This happened because we assumed that the source of the injected electrons has $-\partial f / \partial \epsilon$ profile, and we also projected it onto a $-\partial f / \partial \epsilon$ state. We use this model because we are mainly interested in the decay of the "near ground state" modes of Eq. 8 in the main text, which is exactly proportional to $-\partial f / \partial \epsilon$ state. The angular distribution of scattered particles $\sigma(\theta_i)$, is obtained by setting $\theta_j = 0$:

$$\sigma(\theta_i) = \langle \theta_i | I | 0 \rangle,$$ \hfill (37)

The distribution $\sigma(\theta)$ is shown in Fig. 3 of the main text.

Initial operator $I[\delta(p - k)](p_1)$ has a rotational symmetry in a sense that the integral is only a function of $k = |k|$, $p_1 = |p_1|$, and an angle in-between $p_1$ and $k$. Consequently, the matrix elements $\langle \theta_i | I | \theta_j \rangle$ depend only on the $(\theta_i - \theta_j)$ combination: $\langle \theta_i | I | \theta_j \rangle = G(\theta_i - \theta_j)$. The eigenvalues of such a matrix are readily obtained by applying a discrete Fourier transform to $G(\theta)$. Therefore, the eigenvalues of the operator $\langle \theta_i | I | \theta_j \rangle$ can be obtained in the matrix notation by transforming $\sigma(\theta)$ as

$$\lambda_m = \sum_i e^{-im\theta_i} \sigma(\theta) \Delta \theta_i,$$ \hfill (38)

The quantity $\sigma(\theta_i)$ has the meaning of the transition rate per unit angle, with the dimensionality of $\sec^{-1} \text{rad}^{-1}$. The dependence $\sigma(\theta)$ is constrained by particle conservation

$$\sum_i \Delta \theta_i \sigma(\theta_i) = 0$$ \hfill (39)

and momentum conservation

$$\sum_i \Delta \theta_i \cos \theta_i \sigma(\theta_i) = \sum_i \Delta \theta_i \sin \theta_i \sigma(\theta_i) = 0.$$ \hfill (40)

in two-body collisions. These constraints yield the identities

$$\lambda_0 = \lambda_1 = 0.$$ \hfill (41)

The accuracy with which these relations hold provides a useful check for the precision of our numerical method. As an illustration, the eigenvalue $\lambda_1$ is shown in Fig. 1 of the main text (dashed curve). In general $\sigma(\theta)$ is a sign-changing function, with $\sigma(\theta) < 0$ corresponding to the emission of holes in the backward direction. This behavior is illustrated in Fig. 3 of the main text.

On the side, because of the rotational symmetry we do not need to compute all the entries of the matrix $\langle p_m, \theta_n | I | p_{m'}, \theta_{n'} \rangle$. For our purpose it is sufficient to evaluate the vector $\langle p_m, \theta_n | I | p_{m'}, 0 \rangle$, a quantity that for a rotationally invariant problem contains all the information about the operator and its eigenvalues. Therefore, the computation complexity of the problem is reduced from being quadratic in the number of angular points $N$ to that linear in $N$, while remaining quadratic in the number of radial points $M$. 

Separating contributions of the backscattering and forward scattering processes

In this subsection we describe the method of separation of the backscattering from more noisy forward scattering and show that the details of forward scattering has no significant impact on the backscattering. We use this method to show that the potential numerical problems of our approach described above do not affect the qualitative features of the back-scattering and the eigenvalue hierarchy of odd and even harmonics. The initial assumption about the absolute value of the scattering matrix element in $|V| = \text{const}$. Due to computational problems with precision of the forward scattering, we wish to study a modified potential that would prevent particles from scattering in a forward direction while keeping the backscattering effect intact. To accomplish this we choose a new 2-particle scattering potential that would satisfy $|V| \approx 0$ in the vicinity of $p_1 \approx p'_1$ and $p_1 \approx p'_2$ points. An example of a function that approaches zero when some $p_1 \approx p'_1$ is

$$g(p_1 - p'_1) = \left(1 - \exp\left[-\frac{(p_1 - p'_1)^2}{a^2}\right]\right).$$

We consider 4 values of $a = 1, 1/2, 1/4, 1/8$ to study eigenvalue dependence on the cutoff parameter $a$. To make the scattering matrix element to be explicitly symmetric under the time reversal symmetry and particle permutations, we construct it in a following way:

$$w(p_1, p_2, p'_1, p'_2) \sim |V|^2 g(p_1 - p'_1) g(p_1 - p'_2) g(p_2 - p'_1) g(p_2 - p'_2)$$

instead of $w(p_1, p_2, p'_1, p'_2) \sim |V|^2$. We use the properly normalized scattering probability element from (43) with other parameters including temperature, integration mesh, and numerical integration precision being the same. This calculation repeats all the steps of the main calculation, but takes the transition element to approach zero in the case of forward scattering, which effectively turns it off and allows to omit the singularity during numerical integration of Eq. 21 of the main text.

The plot of backscattering $\sigma(\theta)$ analogous to Fig. 3 of the main text is shown in Fig. 5. The backscattering with separated forward scattering in Fig. 5 resembles the same qualitative properties as the backscattering in Fig. 3 of the main text while showing better numerical results and more abrupt regime change. The distribution enters the low-temperature scaling regime faster: the angular distribution already reaches $T^2$ scaling in the amplitude at the temperature $T = 0.32 T_F$, which can be observed at Fig. 5.

The deviation in behavior of the eigenvalues of odd $m$ harmonics from the behavior of eigenvalues of even $m$ harmonics shows up at the temperatures lower then $T = 0.32 T_F$, its behavior can be seen in Fig. 6. Besides the absence of the forward scattering, such calculation produced result analogous to the main results, which means that the forward scattering plays little role in creating the hierarchy of eigenvalues.

FIG. 4. The mesh in momentum space used in the calculation. The mesh density is nonuniform to achieve better coverage of angles for the collinear and anti-collinear directions relative to the incoming momentum \( \mathbf{k} \). This choice guarantees that the mesh respects rotational symmetry of the problem. In the actual calculation we used \( M = 40 \) radial points, \( N = 200 \) azimuthal points for a less dense mesh (orange points), and \( N' = 200 \) points for a more dense mesh in the collinear and anti-collinear direction of \( \mathbf{k} \) (red points). Accordingly, the total numbers of angles in the collinear and anti-collinear groups was \( N'/2 \); the total number angles in the non-collinear upper and lower groups was \( N/2 \). The radial mesh was chosen to span an annulus covering the Fermi surface (marked by a bold circle). To account for the strong collinear and anti-collinear contributions in the two-body scattering, the \( x, y \) coordinates are rotated so that the \( x \) axis is aligned with the incoming momentum \( \mathbf{k} \).

FIG. 5. The angular distribution \( \sigma(\theta) \) of scattered particles for different temperatures calculated for a potential that effectively separates forward scattering. The temperatures used in this plot are \( T/T_F = 0.0025, 0.005, 0.01, 0.02, 0.04, 0.08, 0.16, 0.32, 0.64, 1.28 \). The value of the coefficient used in (43) is \( a = 1/2 \).
FIG. 6. Eigenvalues for different harmonics as a function of temperature. Even $m$ eigenvalues show the $T^2$ scaling with temperature already at $T = 0.16T_F$. Odd eigenvalues start to diverge from $T^2$ scaling to faster scaling regimes at temperatures lower than $T = 0.32T_F$. The back-scattering shows $T^2$ dependence in the intensity $\sigma$ and $T$ dependence of the width of the scattering peak.