Keldysh Space Control of Charge Dynamics in a Strongly Driven Mott Insulator

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The fate of a Mott insulator under strong low frequency optical driving conditions is a fundamental problem in quantum many-body dynamics. Using ultrafast broadband optical spectroscopy, we measured the transient electronic structure and charge dynamics of an off-resonantly pumped Mott insulator Ca2RuO4. We observe coherent bandwidth renormalization and nonlinear doublon-holon pair production occurring in rapid succession within a sub-100-fs pump pulse duration. By sweeping the electric field amplitude, we demonstrate continuous bandwidth tuning and a Keldysh crossover from a multiphoton absorption to quantum tunneling dominated pair production regime. Our results provide a procedure to control coherent and nonlinear heating processes in Mott insulators, facilitating the discovery of novel out-of-equilibrium phenomena in strongly correlated systems.

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The response of a Mott insulator to a strong electric field is a fundamental question in the study of nonequilibrium correlated many-body systems [1–15]. In the dc limit, a breakdown of the insulating state occurs when the field strength exceeds the threshold for producing pairs of doubly occupied (doublon) and empty (holon) sites by quantum tunneling, in analogy to the Schwinger mechanism for electron-positron pair production out of the vacuum [16]. Recently, the application of strong low frequency ac electric fields has emerged as a potential pathway to induce insulator–metal transitions [17–20], realize efficient high-harmonic generation [21,22], and coherently manipulate band structure and magnetic exchange interactions in Mott insulators [23–28]. Therefore there is growing interest to understand doublon-holon (d-h) pair production and their nonthermal dynamics in the strong field ac regime.

Strong ac field induced d-h pair production has been theoretically studied using Landau-Dykhnje adiabatic perturbation theory [29] along with a suite of nonequilibrium numerical techniques [17,21,22,29–32]. Notably, d-h pairs are primarily produced through two nonlinear mechanisms: multiphoton absorption and quantum tunneling [29,33]. The two regimes are characterized by distinct electric field scaling laws and momentum space distributions of d-h pairs. By tuning the Keldysh adiabaticity parameter $\gamma_K = \hbar \omega_{pump}/(eE_{pump}\xi)$ through unity, where $\omega_{pump}$ is the pump frequency, $E_{pump}$ is the pump electric field, $e$ is electron charge, and $\xi$ is the d-h correlation length, a crossover from a multiphoton dominated ($\gamma_K > 1$) to a tunneling dominated ($\gamma_K < 1$) regime can in principle be induced. However, direct experimental tests are lacking owing to the challenging need to combine strong tunable low frequency pumping fields with sensitive ultrafast probes of nonequilibrium distribution functions.

We devise a protocol to study these predicted phenomena using ultrafast broadband optical spectroscopy. As a test bed, we selected the multiband Mott insulator Ca2RuO4. Below a metal-to-insulator transition temperature $T_{MIT} = 357$ K, a Mott gap ($\Delta = 0.6$ eV) opens within its 2/3-filled Ru 4d $t_{2g}$ manifold [34–37], with a concomitant distortion of the lattice [38]. Upon further cooling, the material undergoes an antiferromagnetic transition at $T_N = 113$ K into a Néel ordered state. It has recently been shown that for temperatures below $T_{MIT}$, reentry into a metallic phase can be induced by a remarkably weak dc electric field of order 100 V/cm [39], making Ca2RuO4 a promising candidate for exhibiting efficient nonlinear pair production.

To estimate the response of Ca2RuO4 to a low frequency ac electric field, we calculated the d-h pair production rate (Γ) over the Keldysh parameter space using a Landau-Dykhnje method developed by Oka [29]. Experimentally determined values of the Hubbard model parameters for Ca2RuO4 were used as inputs [40]. As shown in Fig. 1(a), Γ is a generally increasing function of $E_{pump}$ and $\hbar \omega_{pump}$. For a
fixed $\omega_{\text{pump}}$, the predicted scaling of $\Gamma$ with $E_{\text{pump}}$ is clearly different on either side of the Keldysh crossover line ($\gamma_K = 1$), evolving from power law behavior $\Gamma \propto (E_{\text{pump}})^{\alpha}$ in the multiphoton regime to threshold behavior $\Gamma \propto \exp(-b/E_{\text{pump}})$ in the tunneling regime [Fig. 1(b)].

At time delays where coherent nonlinear processes are absent, the transient pump-induced change in reflectivity of a general gapped material is proportional to the density of photoexcited quasiparticles [51–53], which, upon dividing by a constant pump pulse duration ($\sim$100 fs), yields $\Gamma$. Differential reflectivity ($\Delta R/R$) transients from Ca$_2$RuO$_4$ single crystals were measured at $T = 80$ K using several different subgap pump photon energies ($\hbar \omega_{\text{pump}} < \Delta$) in the midinfrared region, and across an extensive range of probe photon energies ($\hbar \omega_{\text{probe}}$) in the near-infrared region spanning both the $\alpha$ and $\beta$ absorption peaks [Fig. 1(c)]. These two band edge features can be assigned to optical transitions within the Ru $t_{2g}$ manifold [37,54]. Figure 1(d) shows reflectivity transients at various fluences measured using $\hbar \omega_{\text{pump}} = 0.3$ eV and $\hbar \omega_{\text{probe}} = 1.77$ eV. Upon pump excitation, we observe a rapid resolution-limited drop in $\Delta R/R$. With increasing fluence, the minimum value of $\Delta R/R$ becomes larger, indicating a higher value of $\Gamma$ within the pump pulse duration. This is followed by exponential recovery as the $d$-$h$ pairs thermalize and recombine [40]. By plotting $\Gamma$ against the peak value of $E_{\text{pump}}$ (measured in vacuum), we observe a change from power law scaling to threshold behavior when $E_{\text{pump}} > 0.07$ V/Å [Fig. 1(e)], in remarkable agreement with our calculated Keldysh crossover [Figs. 1(a) and 1(b)]. In contrast, measurements performed using 0.56 eV pumping exhibit exclusively power law scaling over the same $E_{\text{pump}}$ range [Fig. 1(f)], again consistent with our model.

A predicted hallmark of the Keldysh crossover is a change in width of the nonthermal distribution of $d$-$h$ pairs in momentum space [29]. In the multiphoton regime, doublons and holons primarily occupy the conduction and valence band edges, respectively, resulting in a pair distribution function ($P_p$) sharply peaked about zero momentum ($p = 0$). In the tunneling regime, the peak drastically broadens, reflecting the increased spatial localization of $d$-$h$ pairs. Using the Landau-Dykhnne method [40], we calculated the evolution of $P_p$ for Ca$_2$RuO$_4$ as a function of $E_{\text{pump}}$ through the Keldysh crossover. Figure 2(a) displays $P_p$ curves at three successively larger $E_{\text{pump}}$ values corresponding to (i) $\gamma_K = 1.49$, (ii) $\gamma_K = 0.75$, and (iii) $\gamma_K = 0.47$, which show a clearly broadening width along with increasing amplitude.

To demonstrate how signatures of a changing $P_p$ width are borne out in experiments, we simulate the effects of different nonthermal electronic distribution functions on the broadband optical response of a model insulator. Assuming a direct-gap quasi-two-dimensional insulator with cosine band dispersion in the momentum plane ($p_x$, $p_y$), the optical susceptibility computed using the density matrix formalism can be expressed as [40,55].
\[ \chi = \sum_{p_x, p_y} C \mathcal{L} \left( \hbar \omega_{\text{probe}} - \Delta(p_x, p_y) \right) \left[ N_e(p_x, p_y) - N_c(p_x, p_y) \right], \]

where \( C \) is a constant incorporating the transition matrix element, \( \mathcal{L} \) represents a Lorentzian oscillator centered at the gap energy \( \Delta(p_x, p_y) \), and \( N_e \) and \( N_c \) are the occupations of the valence and conduction bands, respectively. As will be shown later [Fig. 3(a)], it is valid to assume that \( \Delta(p_x, p_y) \) decreases in proportion to the number of excitations [40]. Figure 2(b) shows simulated reflectivity spectra around the band edge—converted from \( \chi \) via the Fresnel equations—using Gaussian functions for \( N_e \) and \( N_c \) of variable width to approximate the \( P_p \) line shapes [Fig. 2(a)] [40]. As \( P_p \) evolves from condition (i) to (iii), we find that the intersection between the nonequilibrium and equilibrium reflectivity spectra shifts to progressively higher energy. For comparison, we also performed simulations under resonant photodoping conditions using the direct-gap insulator model. Figure 2(c) displays three \( P_p \) curves at successively larger \( E_{\text{pump}} \) values, which were chosen such that the total number of excitations match those in Fig. 2(a). Each curve exhibits maxima at nonzero momenta where \( \hbar \omega_{\text{pump}} = \Delta(|p|) \) is satisfied. In stark contrast to the subgap pumping case, the amplitude of \( P_p \) increases with \( E_{\text{pump}} \) but the width remains unchanged. This results in the nonequilibrium reflectivity spectra all intersecting the equilibrium spectrum at the same energy, forming an isosbestic point [Fig. 2(d)].

The presence or absence of an isosbestic point is therefore a key distinguishing feature between Keldysh space tuning and photodoping. This criterion can be derived from a more general analytical model [40], which shows that a key condition for identifying a Keldysh crossover is that \( \Delta R/R \) spectra at difference fluences do not scale. 

Probe photon energy-resolved \( \Delta R/R \) maps of \( \text{Ca}_2\text{RuO}_4 \) were measured in both the Keldysh tuning (\( \hbar \omega_{\text{pump}} = 0.3 \) eV) and photodoping (\( \hbar \omega_{\text{pump}} = 1 \) eV) regimes. As shown in Figs. 2(e) and 2(f), the extremum in \( \Delta R/R \), denoting the peak \( d-h \) density, occurs near a time \( t = 0.1 \) ps measured with respect to when the pump and probe pulses are exactly overlapped (\( t = 0 \)). This is followed by a rapid thermalization of \( d-h \) pairs as indicated by the fast exponential relaxation in \( \Delta R/R \), which will be discussed later [40]. Figure 2(g) shows \( \Delta R/R \) maps acquired in the subgap pumping regime for three different pump fluences corresponding to conditions (i)–(iii) in Figs. 1(e) and 2(a). Focusing on the narrow time window around \( t = 0.1 \) ps, where the \( d-h \) distribution is highly nonthermal, we observe that \( \Delta R/R \) changes sign across a well-defined probe energy (dashed red line), marking a crossing point of the transient and equilibrium reflectivity spectra. As \( \gamma_K \) decreases, the crossing energy increases, evidencing an absence of an isosbestic point. Analogous maps acquired in the photodoping regime [Fig. 2(h)] also exhibit a sign change. However, the crossing energy remains constant over an order of magnitude change in fluence, consistent with an
that the electronic subsystem indeed thermalizes by rise to very different nonthermal distributions (Fig. 2). Linear and nonlinear pair production processes initially give density. As shown in Fig. 3(b), the system has lost memory of (fluence 26 mJ/cm²). Conversely, by t = 0.5 ps, the curves overlap very well [Fig. 3(c)], indicating that the system has lost memory of isosbestic point. These measurements corroborate our simulations and highlight the unique distribution control afforded by Keldysh tuning.

To study the d-h thermalization dynamics in more detail, we used a Kramers-Kronig transformation to convert our differential reflectivity spectra into differential conductivity (σ) spectra [40]. Figure 3(a) shows the real part of the transient conductivity measured in the thermalized state (t = 0.5 ps) following an 0.3 eV pump pulse of fluence 26 mJ/cm² (τₖ = 0.5), overlaid with the equilibrium conductivity. Subgap pumping induces a spectral weight transfer from the β to α peak and a slight redshift of the band edge, likely due to free carrier screening of the Coulomb interactions [56]. Unlike in the dc limit, there is no sign of Mott gap collapse despite Epump exceeding 10⁹ V/m. To verify that the electronic subsystem indeed thermalizes by t = 0.5 ps, we compare the real parts of Δσ₁ eV (fluence 26 mJ/cm²) and Δσ₂ eV (fluence 4 mJ/cm²), the change in conductivity induced by subgap and above-gap pumping, respectively, at both t = 0.1 and 0.5 ps. A scaling factor A is applied to Δσ₁ eV to account for any differences in excitation density. As shown in Fig. 3(b), the t = 0.1 ps curves do not agree within any scale factor. This is expected because the linear and nonlinear pair production processes initially give rise to very different nonthermal distributions (Fig. 2). Conversely, by t = 0.5 ps, the curves overlap very well [Fig. 3(c)], indicating that the system has lost memory of how the d-h pairs were produced and is thus completely thermalized.

Based on the observations in Figs. 3(b) and 3(c), the nonthermal window can be directly resolved by evaluating the time interval over which the quantity Δ(Δσ) = Δσ₁ eV − A × Δσ₁ eV is nonzero [40]. Figure 4(a) shows the complete temporal mapping of Δ(Δσ) spectra. The signal is finite only around t = 0 ps and is close to zero otherwise, supporting the validity our subtraction protocol. By taking a constant energy cut, we can extract a thermalization time constant of around 0.2 ps [Fig. 4(b)]. Interestingly, ΔR/R and Δσ₀,₃ eV, which both track the d-h pair density, peak near 0.1 ps, whereas Δ(Δσ) peaks earlier at t = 0 when the d-h pair density is still quite low. This implies the existence of an additional coherent
nonthermal process that scales with \( E_{\text{pump}} \), which peaks at \( t = 0 \), rather than with the \( d\!-\!h \) density.

To identify the physical process responsible for the \( t = 0 \) signal, we examined how the electronic structure of \( \text{Ca}_2\text{RuO}_4 \) would need to change in order to produce the \( \Delta(\Delta\sigma) \) profile observed at \( t = 0 \) [Fig. 4(c)]. Using density functional theory (DFT), we performed an \textit{ab initio} calculation of the optical conductivity of \( \text{Ca}_2\text{RuO}_4 \) based on its reported lattice and magnetic structures below \( T_N \). The tilt angle of the \( \text{RuO}_6 \) octahedra was then systematically varied in our calculation as a means to simulate a changing electronic bandwidth [40]. We find that both the real and imaginary parts of the measured \( \Delta(\Delta\sigma) \) spectrum at \( t = 0 \) are reasonably well reproduced by our calculations if we assume the bandwidth of the driven system \( (W) \) to exceed that in equilibrium \( W_{\text{eq}} \) [Fig. 4(d)] [40]. This points to the coherent nonthermal process being a unidirectional ultrafast bandwidth renormalization (UBR) process that predominantly occurs under subgap pumping conditions [40].

Coherent UBR can in principle occur via photoassisted virtual hopping between lattice sites, which has recently been proposed as a pathway to dynamically engineer the electronic and magnetic properties of Mott insulators [23–28]. To quantitatively extract the time and \( E_{\text{pump}} \) dependence of the fractional bandwidth change \( (W - W_{\text{eq}})/W_{\text{eq}} \), we collected \( \Delta(\Delta\sigma) \) spectra as a function of both time delay and pump fluence and fit them to DFT simulations [40]. As shown in Fig. 4(e), the bandwidth change exhibits a pulse-width limited rise with a maximum \( t = 0 \) value that increases monotonically with the peak pump field, reaching up to a relatively large amplitude of 1.5\% at \( E_{\text{pump}} = 0.12 \text{ V/Å} \), comparable to the bandwidth increases induced by doping [36] and pressure [57]. Independently, we also calculated the field dependence of \( (W - W_{\text{eq}})/W_{\text{eq}} \) expected from photoassisted virtual hopping by solving a periodically driven two-site Hubbard model in the Floquet formalism [23,40], using the same model parameters for \( \text{Ca}_2\text{RuO}_4 \) as in our Landau-Dykhnne calculations [Fig. 1(a)]. We find a remarkable match to the data without any adjustable parameters [Fig. 4(e)]. Since bandwidth renormalization increases with the Floquet parameter \( (eaE_{\text{pump}})/\hbar\omega_{\text{pump}} \) in the case of photoassisted virtual hopping, where \( a \) is the intersite distance, this naturally explains why subgap pumping induces the much larger UBR effect compared to above-gap pumping.

The ability to rationally tune a Mott insulator \textit{in situ} over Keldysh space enables targeted searches for exotic out-of-equilibrium phenomena such as strong correlation assisted high-harmonic generation [21,22], coherent dressing of quasiparticles [58], Wannier-Stark localization [2,17], ac dielectric breakdown [29], and dynamical Franz-Keldysh effects [32,59], which are predicted to manifest in separate regions of Keldysh space. It also provides control over the nonlinear \( d\!-\!h \) pair production rate—the primary source of heating and decoherence under subgap pumping conditions—in Mott systems, which is crucial for experimentally realizing coherent Floquet engineering of strongly correlated electronic phases.

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