Transition from synchronous to asynchronous superfluid phase slippage in an aperture array

Y. Sato, E. Hoskinson, and R. E. Packard

Department of Physics, University of California, Berkeley, California 94720, USA

(Received 30 August 2006; published 5 October 2006)

We have investigated the dynamics of superfluid phase slippage in an array of apertures. The magnitude of the dissipative phase slips shows that they occur simultaneously in all the apertures when the temperature is near \( T_s \sim T \sim 10 \text{ mK} \) and subsequently lose their simultaneity as the temperature is lowered. We describe three experiments to probe the mechanisms underlying the synchronous behavior. The results raise fundamental questions about the dynamics of phase slippage in a multiply connected geometry.

DOI: 10.1103/PhysRevB.74.144502

PACS number(s): 67.40.Vs, 05.45.Xt, 67.40.Hf, 67.40.Rp

I. INTRODUCTION

Superfluid \(^4\text{He}\) is described by a complex order parameter \( \psi \propto e^{i\phi} \). Phase differences are proportional to superfluid velocity and vary as \( d(\Delta \phi)/dt = -\Delta \mu / \hbar \). Superflow is driven by chemical potential differences, \( \Delta \mu = m_3 (\Delta P / \rho - s \Delta T) \), where \( \Delta P \) and \( \Delta T \) are differences in pressure and temperature, \( \rho \) is the mass density, \( s \) is the specific entropy, and \( m_3 \) is the \(^4\text{He}\) atomic mass.\(^{1,2}\) Whenever the flow through a submicron-size aperture reaches a critical velocity \( \nu_c \), dissipation occurs in a discrete event wherein the quantum phase difference across the aperture drops by \( 2\pi \).\(^{3-5}\) Since superfluid velocity is proportional to phase gradient, this \( 2\pi \) “slip” corresponds to a discrete drop in velocity, \( \nu_{\text{slip}} = \kappa / l_{\text{eff}} \) where \( \kappa = \hbar / m_3 \) is the quantum of circulation and \( l_{\text{eff}} \) is the effective hydrodynamic length of the aperture.

If one applies a constant chemical potential difference \( \Delta \mu \) across an aperture, the superfluid velocity increases linearly to the critical velocity, followed by an abrupt drop (if the duration of the slip is short compared to the relaxation time) and followed again by a linear increase. The waveform of superfluid velocity \( \nu_{\phi}(t) \) then resembles a sawtooth in which the phase slip events take place at an average rate equal to the Josephson frequency \( f_j = \Delta \mu / \hbar \) (Ref. 6). For single apertures, stochastic fluctuations in the critical velocity usually obscure the periodic nature of this process.\(^{7}\)

Recent work\(^8\) has shown that in superfluid \(^4\text{He}\) periodic phase slip oscillations at frequency \( f_j \) exist in an array of \( N \) (\( \sim 4225 \) ) apertures. The oscillation amplitude near the superfluid transition temperature implies that the phase slips occur synchronously (i.e., simultaneously) among all the \( N \) apertures. Josephson oscillations can be used as a phase difference sensor in superfluid gyrosopes and interferometers.\(^9-11\)

It is necessary to understand the origin of the synchronicity mechanism in order to optimize the design of such devices.

To investigate the nature of phase slips within the array, we have performed three kinds of experiments. In the first, we drive phase slip oscillations by applying a chemical potential difference across an aperture array and measure the phase slip oscillation amplitude down to \( T_s \sim T \sim 160 \text{ mK} \). We find that the amplitude decreases rather dramatically as the temperature is lowered, as compared to what would be expected for synchronous behavior. In a second experiment, we excite transient Josephson oscillations lasting from one cycle to thousands of cycles. We find that the phase slip size does not change over many cycles of oscillation, indicating that when phase slips are synchronous, they are synchronous from the very first slip. In the third experiment, we give the system an initial excitation energy, allow it to decay through the dissipative phase slips, then record the amplitude of the subcritical current oscillation (the so-called Helmholtz mode) that occurs after the last phase slip. We find that as the temperature decreases, phase slips within the array seem to occur in a less abrupt manner implying that a phase slip event is no longer a single simultaneous array-wide event but rather a collection of uncorrelated events localized to individual apertures. We present these three findings in the first part of this paper and discuss possible interpretations in the second.

II. TYPE I EXPERIMENT

Our experimental apparatus is shown in Fig. 1. Two volumes filled with superfluid \(^4\text{He}\) are separated by a diaphragm and an array of \( N = (4225) \) apertures that are \( \sim 30 \text{ nm} \) in diameter and spaced \( 3 \mu \text{m} \) apart in a \( 50 \text{ nm} \) thick silicon nitride chip. A thin, flexible, metal-coated diaphragm can be pulled toward an electrode by the application of a voltage between them. A SQUID-based displacement sensor\(^12\) is used to monitor the position of the diaphragm that serves as a microphone to determine the magnitude of the phase slip oscillation.

In our first type of measurement, we apply a DC step voltage between the diaphragm and the electrode at \( t = 0 \).

This pulls the flexible diaphragm toward the electrode creating a pressure head (and therefore a chemical potential difference) across the array. If the initial pull is large enough, the flow velocity inside the apertures reaches \(v_t\), and the fluid undergoes \(2\pi\) phase slips at the Josephson frequency. These dissipative events continue until there is no energy left to drive the fluid up to the critical velocity. The phase slip oscillations end, and the system begins to oscillate about \(\Delta \mu = 0\) at a different frequency—the Helmholtz frequency. The restoring force of the diaphragm, the inertia of the fluid moving in the apertures, and the heat capacity of the fluid in the inner volume determine the frequency of this resonant mode. Figure 2(a) shows a typical diaphragm displacement \(x(t)\) during one of these relaxation transients. The discontinuities in fluid velocity due to phase slip events show up as sudden slope changes in \(x(t)\). These can be seen in the first half of Fig. 2(b).

To determine whether or not phase slips are occurring synchronously throughout the array, we measure the peak-to-peak amplitude of the phase slip current oscillations, \(I_{\text{slip}}\), and compare this number to the expected magnitude if all \(N\) apertures are locked together, \(I_{\text{slip}}^N\). This expected magnitude is determined by directly measuring the current phase relation \(I(\phi)\) for the array during periods of subcritical flow (i.e., the Helmholtz oscillation) where the flow is synchronous across the array. We are concerned with the strong coupling regime \(T_\chi - T \approx 10\) mK, where \(I(\phi)\) is linear. The expected magnitude of the current oscillation for synchronous \(2\pi\) phase slips is then

\[
I = \rho A x, \tag{2}
\]

where \(\rho\) is the total fluid density, and \(A\) is the diaphragm area.

When a chemical potential differential exits across the array, the diaphragm exhibits oscillations at the Josephson period, \(f_j^{-1}\). If the amplitude of such diaphragm oscillations is \(x_a\), the magnitude of the mass current oscillations at frequency \(f_j\) is given by

\[
I_{\text{slip}} = \frac{2\pi f_j \rho A x_a}{\gamma}, \tag{3}
\]

where \(\gamma\) is the Fourier coefficient of the first harmonic of the displacement sensor signal. We assume here that the current exhibits a sawtooth waveform, a case where \(\gamma = 2/\pi\).

To determine \(x_a\), we record the signal \(x(t)\) preceding the Helmholtz mode and compute the Fourier transform of the diaphragm oscillations. By analyzing the spectral content in small time intervals, we extract the frequency and the amplitude of the phase slip oscillations as a function of time throughout the transient. Once we obtain the amplitude of the diaphragm oscillations \(x_a\), we use Eq. (3) to compute \(I_{\text{slip}}\). The mass current oscillation amplitude \(I_{\text{slip}}\) varies with frequency due to cell resonances but levels off at lower frequencies (typically below 300 Hz). We use this limiting value for \(I_{\text{slip}}\). An example of this frequency dependence of \(I_{\text{slip}}\) is shown in Fig. 3.

The Fourier analysis becomes more difficult at lower temperatures because the duration of the phase slip flow becomes shorter due to increasing critical velocity. To extend the duration of phase slip flow, we use a heater installed...
inside of the inner cell. First, we apply a step voltage to the heater which creates a temperature differential $\Delta T$ across the array and starts the phase slip oscillation. We then continuously increase the heater power during the transient to counteract cooling due to net superfluid flow through the array (the thermomechanical effect). In this way, we slow the rate at which the chemical potential goes to zero. The extended transient allows us to apply the Fourier analysis described above and find the amplitude of oscillations at lower temperatures.

Figure 4 shows the variation of $I_{\text{slip}}$ with temperature. For comparison, we also plot $I_{\text{slip}}^N$ defined by Eq. (1) using data derived from Ref. 14.

As seen in the figure, at the highest temperatures where the phase slips appear, $(T_h - T = 9 \text{ mK})$, we find $I_{\text{slip}} = I_{\text{slip}}^N$, which implies that phase slips are occurring synchronously among all the $N$ apertures. However, as the temperature decreases, the amplitude of current oscillation starts to rapidly decrease (relative to $I_{\text{slip}}^N$) showing a loss of synchronicity among apertures. This is the central finding of this experiment.

### III. TYPE 2 EXPERIMENT

Neither the mechanism for the initial synchronization nor the reason for its subsequent loss is yet fully understood. However, systems of interacting nonlinear oscillators often exhibit synchronization after multiple cycles. If such nonlinear mode locking is present in the array, one would expect the size of first phase slip to be smaller than that of the $n^{th}$ where $n \gg 1$. Our second type of experiment is directed toward determining if there is a change in overall slip size between the first and nth phase slip oscillation.

Equation (2) shows that when a dissipative phase slip occurs, the sudden current drop in the aperture array is reflected by a sudden change in the slope of the diaphragm position curve $x(t)$. By adjusting the voltage step applied to the diaphragm we vary the length of the phase slip oscillation train from as little as one slip to as many as several thousands of slips. We then compare the abrupt slope changes, shown in Fig. 2(b), at the first slip and the $n^{th}$ slip.

![FIG. 4. Measured phase slip current oscillation amplitude $I_{\text{slip}}$ (for $f < 300 \text{ Hz}$) and the expected value for a fully synchronous case $I_{\text{slip}}^N$. The lines are a guide to the eye.](image)

The change in the slope, $\Delta x(t)$ is determined as follows. The fluid acceleration is proportional to the chemical potential difference $\Delta \mu$ across the array. If $\Delta \mu$ is constant in the vicinity of a slip, the current increases linearly in time and the displacement of the diaphragm follows a parabola. We fit two parabolas at the cusp in the diaphragm position $x(t)$ (one before the phase slip and another right after) and find the change in the slope $\Delta x(t)$. An example of this parabolic fit is shown in Fig. 5.

The measured slope changes at the first slip and the $n^{th}$ slip are plotted in Fig. 6. We find that the phase slip size does not change over many cycles. This result shows that when the oscillations are synchronous, they are synchronous from the very first slip. We conclude then that the synchronization is not due to a typical nonlinear mode locking process.

### IV. TYPE 3 EXPERIMENT

Our third experiment sheds additional light on the nature of collective phase slippage in the array. We apply a small step voltage, $V$, between the diaphragm and the electrode to create chemical potential differentials which are sufficiently

![FIG. 5. Typical parabolic fits. The cusp shown is the last phase slip before the Helmholtz oscillation in Fig. 2(b). Dots are the data, and solid lines are the fits. We fit two parabolas at the cusp and find the change in the slope.](image)

![FIG. 6. Diaphragm velocity change at the first and the $n^{th}$ slip where $n$ is on the order of 1000. The temperature dependence comes from the increasing superfluid density as the temperature decreases.](image)
small to keep the fluid velocity inside the apertures subcritical. In the subsequent flow transient, the chemical potential reaches zero without inducing any phase slips and the diaphragm oscillates at the Helmholtz frequency with an initial amplitude \( x_0 \). In the absence of phase slippage, the initial energy in the Helmholtz oscillation, \( E_h \), should be proportional to \( E_0 \), which is the energy that we put into the system by the application of a voltage step. As we increase the initial kick on the diaphragm and plot \( E_h \) versus \( E_0 \), we expect a line with constant slope until \( E_0 \) is large enough to accelerate the fluid up to \( \nu_c \), triggering a phase slip. At that point, energy is dissipated. If phase slips occur simultaneously in all the \( N \) apertures, \( E_h \) should then drop discontinuously due to the abrupt extraction of energy. After such an event, as we increase \( E_0 \) further, \( E_h \) should increase linearly again until the process repeats.

Since the equilibrium diaphragm displacement is proportional to \( V^2 \), the energy that we put into the system, \( E_0 \), scales as \( V^4 \). The initial energy in the Helmholtz oscillation, \( E_h \), is proportional to the square of the initial Helmholtz diaphragm oscillation amplitude, \( x_0^2 \). Thus a plot of \( x_0^2 \) versus \( V^4 \) (which corresponds to \( E_h \) versus \( E_0 \)) should be a sawtooth if the phase slippage occurs abruptly and simultaneously throughout the array. If the phase slippage process is distributed in time, as individual apertures slip independently of others, the sawtooth would be rounded.

Figure 7 shows our measurements of \( x_0^2 \) vs \( V^4 \) at various temperatures. As the temperature is lowered below \( T_A \), the shape of \( x_0^2 \) vs \( V^4 \) evolves from a sharp sawtooth indicative of an abrupt collective phase slip event to a smoother curve that implies a continuous “phase slide” process. This suggests that some apertures are experiencing a phase slip before the others, allowing the array to dissipate energy in a more continuous manner.

Figure 7 also illustrates the striking crossover from a dissipative phase slip regime to the nondissipative Josephson regime. The critical velocity \( \nu_c \) (or Helmholtz amplitude) at which a slip occurs increases as the temperature decreases. At \( T_A - T = 15 \) mK, \( \nu_c / \nu_{slip} \) and a single array phase slip event removes almost all the energy in the fluid and leaves none for the Helmholtz mode. Therefore, the Helmholtz oscillation amplitude goes to 0 every time a phase slip occurs. As one gets closer to \( T_A \), \( \nu_c \) becomes smaller than \( \nu_{slip} \), and a phase slip event causes a reversal in the flow direction. Phase slips are no longer fully dissipative—the system retrieves some of the energy involved in the reversal of flow. At \( T_A - T = 5 \) mK, where \( \nu_{slip} \approx 2 \nu_c \), dissipation due to the oscillations, which are still present as Josephson oscillations instead of phase slips, ceases. One can view this to be the complete transition into a weakly coupled Josephson regime. In the weakly coupled regime the dominant dissipation occurs through thermal conduction and normal flow—Josephson oscillations cease and Helmholtz oscillations begin when there is no longer enough energy (the flat limiting value in the 5 mK data) to reach the critical current and drive Josephson oscillations. This alternate form of dissipation, although small compared to the phase slips, explains why (in the phase slip regime) the period of the \( x_0^2 \) versus \( V^4 \) curves increases with \( V^4 \); for larger initial energy, the system takes longer to reach the Helmholtz mode and more energy is dissipated through thermal conduction and normal flow.

**V. INTERPRETATIONS**

We have considered possible mechanisms for the observed decrease in phase slip amplitude as exhibited in Fig. 4. Discrete phase slippage in superfluid \(^4\)He is usually associated with the passage of quantized vortices that are stochastically nucleated near the aperture surface. The intrinsic fluctuations cause the critical velocity to be spread out over a range \( \Delta \nu_c \). This finite distribution width can cause the phase slip oscillation to lose its well-defined periodicity.

The critical velocity width \( \Delta \nu_c \) is a function of temperature, and the relevant quantity in determining the temporal coherence of phase slip oscillations in a given aperture is \( \Delta \nu_c / \nu_{slip} \). If \( \Delta \nu_c / \nu_{slip} > 1 \), the periodicity at \( f_1 \) is lost. Previous work\(^7,21,22\) suggests that this ratio \( \Delta \nu_c / \nu_{slip} \) increases with decreasing temperature near the superfluid transition temperature. The observed decline in the oscillation amplitude therefore could be a manifestation of loss of periodicity in any individual aperture.

Another possible mechanism for the loss of synchronicity at lower temperatures may involve variations in the surface microstructure among the array apertures. With the fluid flowing fastest near asperities, the critical velocity for an aperture must be affected by the surface inhomogeneities. Since the superfluid healing length \( \xi \) is a function of temperature, how much of these nanoscale inhomogeneities the fluid actually “sees” should depend on temperature as well. The healing length is given by

\[
\xi(T) = \frac{0.3 \text{ nm}}{(1 - T/T_A)^{0.67}}
\]

(4)
and it decreases from \(\sim 10\) nm to \(\sim 1.5\) nm as the temperature is lowered from \(T_\lambda - T = 10\) mK to \(T_\lambda - T = 160\) mK. If the surface variations are on the order of a few nanometers, this could very well provide a critical velocity distribution whose width increases with decreasing temperature while allowing the individual apertures to maintain well-defined periodic oscillations.

Several overarching questions remain. Is it possible for apertures to act independently in the presence of a macroscopic wave function? Circulation around every loop drawn through the apertures must be quantized while minimizing the energy associated with the phase gradient across the array. It is not clear how this condition is satisfied when phase slips are occurring in random positions within the array.

What are the dynamics of vortices near the transition temperature when the energy removed in a single phase slip becomes comparable to the flow energy itself? What is even meant by a “vortex” when the vortex core \(\sim \xi(T)\) is comparable to the size of the apertures? Perhaps then phase slips occur by collapse of the wave function rather than by vortex dynamics.\(^{23}\) The superfluid order parameter may already be so weakened that at \(\nu_c\) the fluid in the aperture becomes momentarily normal before superfluidity is restored to a state in which the phase difference across the array has dropped by \(2\pi\). This might lead to synchronicity if the wave function is so weak in all of the apertures that an excitation that causes the wave function to collapse in one aperture perturbs the other apertures enough to cause them all to collapse.

VI. CONCLUSION

The experiments described above show that near \(T_\lambda\) phase slippage occurs collectively in all the apertures in an array and the related oscillations at the Josephson frequency are not due to nonlinear mode locking. The observed decline in phase slip oscillation amplitude and the rounding of the sawtooth in the \(x_\lambda^2\) versus \(V^2\) plot both indicate that array phase slippage loses its collective nature as the temperature is lowered. The results reported herein raise fundamental questions about the phase slippage process and the nature of a weakened superfluid confined in a multiply connected region.

Based on our experimental results, Pekker et al.\(^{24}\) have recently constructed a model to understand the phase slip dynamics through an aperture array. Their model couples all the apertures through the bulk superfluid and allows one to investigate how the critical velocity distribution among various apertures affects the experimental observables. The results they obtain with a mean-field approximation and exact numerical analysis for a small number of apertures seem to capture the general features of our findings. An important element that is made clear in their theoretical work is “the competition between quenched disorder and interactions.”\(^{25-27}\) Pekker et al. suggest that further investigations with different numbers of apertures, sizes and spacing might reveal a similar transition in superfluid phase slippage in an aperture array.

ACKNOWLEDGMENTS

We thank Aditya Joshi for his suggestions to the manuscript. We acknowledge stimulating conversations with Dung Hai Lee and Henry Fu. We thank David Pekker, Roman Barankov, and Paul Goldbart for sharing their insights and building on our experimental results. This work was supported in part by the NSF Grant No. DMR 0244882 and NASA.

---

2. D. R. Tilley and J. Tilley, *Superfluidity and Superconductivity*, 3rd edition (IOP, New York, 1990), Sec. 3.3.


