Anomalous proximity effect in an inhomogeneous disordered superconductor

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(Dated: March 22, 2002)

By combining very low temperature scanning tunneling microscopy and spectroscopy on a TiN film we have observed a non uniform state comprising of superconducting (S) and normal (N) areas. The local density of states displays a spatial dependence between S and N different from the usual proximity effect. We conclude that mesoscopic fluctuations might play a major role in accordance with recent theories on superconductor-normal metal quantum transition.

PACS numbers: 74.45.+c, 74.81.-g, 71.30.+h

In two dimensions, the superconductor-insulator transition (SIT) is traditionally described by two microscopic mechanisms \([1]\). In the first one, the Cooper pairs are all formed at \(T=0\) and become localized through the SIT. This is often referred to as the “bosonic” model and has been successfully to granular superconductors \([2]\). The onset of the superconductivity appears at a constant temperature but the transition becomes broader as the sheet resistance \(R_{\parallel}\) approaches \(\frac{\hbar}{2e^2}\). This broadening is associated with quantum fluctuations of the phase of the order parameter due to the competition between the charging energy of the superconducting grains and the Josephson coupling between them. The second mechanism attributes the weakening of the superconductivity to a disorder enhanced Coulomb repulsion. Pair breaking is considered simultaneously with the decreasing of both the superconducting critical temperature and the amplitude of the order parameter as the SIT is approached. This so-called “fermionic” scenario usually describes homogeneous thin films which keep a sharp superconducting transition until disorder becomes very close to a critical value \(\delta\). By further increasing the disorder, by reducing the thickness or by application of a magnetic field, these films can be driven into an insulating state with localized electronic excitations. However, real samples may be neither purely bosonic or purely fermionic systems \([3]\) and a unified theory describing these two limiting cases is still needed. Moreover, there are theoretical predictions \([4]\) and experimental observations \([5, 6, 7, 8, 9]\) of an intermediate metallic phase right at the SIT.

In three dimensions, depending on the disorder, both a SIT and a superconductor-normal metal transition (SMT) are predicted at \(T=0\) \([10, 11, 12]\) and have been experimentally observed in a field-tuned transition \([10]\). The existence of an intermediate metallic phase is often accompanied by a re-entrant temperature and field dependence of the electrical resistance \(R\) of the film \([11]\). Actually, the competition between Cooper pairs and quasiparticle tunneling between superconducting grains or clusters can explain a nonmonotonic behavior of \(R\). In these models, a zero resistance state appears when the network of superconducting islands percolate \([12]\). Inhomogeneities in the morphology of granular films \([13]\) or strong statistical disorder in nominally homogeneous films can lead to the formation of these islands \([14]\).

Mesoscopic effects induced by local fluctuations in the density of randomly distributed impurities can also drive the film into a multiple reentrant field-tuned SMT \([15]\).

A double reentrant behavior very close to a SIT has recently been observed in disordered TiN thin films \([16]\). This SIT could be triggered either by a magnetic field or by increasing \(R_{\parallel}\). It is worth noticing that such a double reentrance has also been observed in Josephson junction arrays and highly granular superconductors \([17]\). In this Letter we report spectroscopic measurements on a similar TiN film with a lower \(R_{\parallel}\) and in zero field, thus far away from the above mentioned SIT. Nevertheless, we show that disorder induced inhomogeneities already exist in such a film which cannot be detected by transport measurements.

TiN was prepared by DC reactive magnetron sputtering at 350°C on thermally oxidized Si substrates. By sputtering a Ti target at various nitrogen partial pressure in an argon-nitrogen gas mixture, different TiN\(_{\delta}\) compounds with \(0.7 \leq \delta \leq 1.2\) can be obtained \([18]\). We have been able to vary the room temperature electrical resistivity between 80 \(\mu\Omega\) cm and 1100 \(\mu\Omega\) cm by changing the nitrogen flow rate from 40 to 200 sccm \([19]\). In stoichiometric TiN films, the mean free path \(l\) is mainly limited by the grain boundaries and one gets \(l \approx L_g\), where \(L_g\) is the typical grain size. When the nitrogen flow rate is increased during the film preparation, Ti vacancies are introduced and one obtains overstoichiometric TiN\(_{\delta}\) (\(\delta > 1\)) with \(l < L_g\). Moreover the Ti vacancies are not uniformly distributed inside the grains but are rather concentrated at their boundaries \([20]\).

The film studied here has an intermediate resistivity of 270 \(\mu\Omega\) cm, a mean free path \(l = 5 \pm 1 \, nm\) and a thickness of 100 nm \([19]\). We estimate \(L_g \approx 20 \, nm\) from STM images. It undergoes a sharp superconducting transition at \(T_c = 4.68 \, K\) as detected by \(R(T)\) experiments.

We have combined topography and spectroscopy measurements with a scanning tunneling microscope (STM) cooled down to very low temperature in a dilution refrigerator. In order to probe the local density of states (LDOS), a small AC modulation of 20 \(\mu V\) rms is added.
FIG. 1: Several spectra taken at different temperatures at the same location and the corresponding BCS fits. Inset: temperature dependence of the BCS gap.

to the sample-tip DC bias voltage $V$ and the differential conductance $\frac{dI}{dV}$ is obtained with a lock-in amplifier technique. At any position above the sample $V$ is ramped and the resulting $\frac{dI}{dV}(V)$ curve gives the LDOS around the Fermi energy. Several spectra are displayed in Fig. 1 at different temperatures. The superconducting gap values $\Delta(T)$ are obtained by convoluting a BCS density of states with a Fermi distribution function and are displayed in the inset. A weakly temperature dependent Dynes parameter in the range $\Gamma = 0.022 \text{ mV}$ to $0.027 \text{ mV}$ is used to adjust the peak height [21]. The lowest indicated temperature of 250 mK is not the measured one which was 125 mK but rather the temperature needed to correctly fit the spectrum. This indicates that the energy resolution of our STM is probably limited by unfiltered electromagnetic radiations which heat the electrons. We find $\Delta(T=0) = 0.73 \text{ mV}$ and a ratio $\frac{\Delta}{kT_c} = 1.81$ not far from the theoretical BCS value of 1.76.

However, these spectra are not observed everywhere on the surface of the TiN film and at other locations a normal-metal like flat LDOS is measured. In order to get an image of the superconducting and normal areas we set the bias to a voltage $V_0 = 0.75 \text{ mV}$ slightly above the BCS gap value. Two images were then recorded simultaneously: the topographic one $Z(x,y)$ and the spectroscopic one $\frac{dI}{dV}(x,y)$ at a given energy $eV_0$. When the tip is scanning above a superconducting region, the differential conductance signal increases because of the peak in the LDOS at $V_0$. Inversely, a flat LDOS is characterized by a lower output of the lock-in amplifier. The two images can be merged into a 3D coloured picture as shown in Fig. 2 where the dark orange areas are normal while the light yellow ones are superconducting [22].

Sometimes, isolated grains stand higher on top of the surface. They can be either normal or superconducting as is shown, respectively, in the upper and lower picture of Fig. 2. Once the electronic properties are identified everywhere on the scanned surface, it is then possible to measure the complete spectra along a line which crosses the boundary between a superconducting and a normal region. Several spectra along such a scan are displayed in Fig. 3. Except for the truly gaped one which is labeled as the starting position $x_{\text{exp}} = 0$ on the S side, none of the spectra are BCS-like. The distance $d_{SN}$ over which the LDOS is varying depends on the local granularity. It can be as long as 50 nm when the surface is smooth. But if we scan across an isolated superconducting grain such as those in the bottom panel of Fig. 2, the spatial variation of the LDOS is much more rapid and takes place within less than 10 nm, mainly inside the grain itself. However, these different spatial scales do not modify the form of
The overall transition of the LDOS at the SN interface since the spectra evolve in the same manner between S and N wherever they are obtained on the surface of the sample.

The spatial dependence of the proximity effect at an SN interface can be described in the framework of the quasiclassical Green’s functions by a complex pairing angle \( \theta(E, r) \) [24]. The LDOS \( n \) at a position \( r \) and for an energy \( E \) from the Fermi level is related to \( \theta \) by \( n(E, r) = n_0\text{Re}[\cos \theta(E, r)] \). In the dirty limit, i.e. \( l \ll \xi = \sqrt{\hbar D/2\Delta} \) (\( D \) is the diffusion constant), \( \theta(E, r) \) obeys the Usadel equation which in one dimension can be written as [24]:

\[
\frac{\hbar D}{2} \frac{\partial^2 \theta}{\partial x^2} + \left[ iE - \Gamma_{in} - 2\Gamma_{sf} \cos \theta \right] \sin \theta + \Delta(x) \cos \theta = 0, \quad (1)
\]

where \( \Gamma_{sf} \) and \( \Gamma_{in} \) are the spin-flip and the inelastic scattering rates respectively. This description of the superconducting proximity effect at a mesoscopic scale has already been checked experimentally in SN heterojunctions with nanofabricated tunnel junctions and with STM experiments [25]. For infinite SN systems, it has been shown that the LDOS exhibits a V-shaped pseudo-gap with peaks below and above the Fermi level. These peaks are separated by an energy which decreases as a function of the distance from the interface in the normal side of the junction. Here, we observe instead U-shaped spectra with peaks that are pinned at the BCS energy \( \Delta \) for any position between S and N. The LDOS is flat for energies smaller than \( \Delta \) and increases as the tip is moved progressively away from the S region. We tried to fit these results using Eq. (1) and a self-consistently determined space dependent order parameter \( \Delta(x) \) [21]. However, no set of parameters was able to correctly reproduce even qualitatively the shape of the spectra. One of the possible reasons for this failure could be that our film does not fulfill the dirty limit condition. Actually, we have \( \xi = 6.5 \text{ nm} \) not much bigger than \( l \) [11]. Moreover, since the scattering centers are concentrated at the grain boundaries, the electronic trajectories could be considered to be quasiballistic inside the grains. The projection in only one dimension of the Usadel equation could also be a too crude approximation.

It is nevertheless striking than we can reproduce with a good accuracy our results if we assume the pairing angle varies linearly with distance:

\[
\frac{\partial \theta}{\partial x} = \frac{\theta_{BCS} - \theta_N}{d_{SN}} \quad (2)
\]

where \( \tan(\theta_{BCS}) = \frac{\Delta}{E+\Delta} \) and \( \theta_N = 0 \). Eq. (2) has been integrated with the boundary conditions,

\[
\theta(x = 0) = \theta_{BCS} \quad \text{and} \quad \theta(x = d_{SN}) = 0 \quad (3)
\]

The numerical results are shown in Fig. 3. We found an excellent agreement between the experimental data and the calculations without any adjustable parameters. \( \Delta \) and \( \Gamma \) are deduced from the spectrum taken above the BCS superconducting region and positions are chosen in order to match the measured LDOS at the Fermi energy. The slight differences between these positions and the measured ones reflect the granularity always present; the latter affects the measured value of the proximity length \( d_{SN} \) and can also perturb the pure linear behavior of Eq. (2). Although, there is no obvious physical meaning behind this equation, we want to point out the universal character of this observed anomalous proximity effect since it does not depend on any physical properties of the material such as inelastic scattering times, conductivities or coherence lengths of S and N necessary to describe the usual proximity effect [27].

We want to discuss now the physical origin of the superconducting and normal clusters. In STM experiments, surface contamination must be considered seriously as a possible artefact for samples exposed to air. Nevertheless, there are easily recognizable signs which indicate the presence of adsorbates: (i) they usually degrade the quality of the images. (ii) At moderately low temperature, when they are frozen out, their positions are revealed as bumps in the topographic images. (iii) At very low temperature, they generally exhibit the LDOS of an insulator and give dark areas in spectroscopic images. As seen in the pictures of Fig. 2, no such effects are visible. The transition between superconducting and normal areas is in places very smooth with no change in the noise in the data. This proves that the inhomogeneities are intrinsic to the sample. However, we cannot rule out that because of oxidation of intergrain boudaries, for example, the surface of TiN itself is not representative of the
bulk which could be more homogeneous. Another possible source of inhomogeneities, could come from spatial variations of the nitrogen concentration in TiN, since the BCS coupling constant, $\lambda$, depends on $\delta$. Following Ioffe and Larkin's pioneering work, strong statistical fluctuations of $\lambda(r)$ can lead to an inhomogeneous system where superconductivity appears first in localized drops which percolate at $T_c$ even far away from the SIT threshold $T_s$. A spatial distribution of the order parameter amplitude would give a broadened density of states as obtained by Hsu et al with large area tunnel junctions.

Here, we would expect instead to observe well identified superconducting clusters with different BCS gap values. It seems therefore more likely that our film is made of well coupled superconducting grains with a unique $\Delta$ embedded in a normal metal matrix where the Ti vacancies pile up. According to Feigel'man et al and Spivak et al, quantum fluctuations in such a system make superconductivity unstable. Spatial variations of the size and the concentration of grains as well as mesoscopic fluctuations of the intergran conductance can drive the film locally into either a normal metal or a superconducting cluster. This is also consistent with the observed leveling off of the resistivity at very low temperature and with the reentrant field tuned transition observed in thinner TiN films. Indeed, when the film is closer to the percolation threshold of the superconducting network its global resistivity is then governed by very few bottlenecks and the associated mesoscopic fluctuations. According to Spivak et al, these mesoscopic fluctuations can give a multiple reentrant transition between superconducting and normal metallic states.

In summary, we have observed inhomogeneities in a disordered superconducting thin film of a nature similar to those invoked to explain macroscopic properties and possibly described by recent fluctuations driven SMT models. It is very likely that in order to understand the spatial dependence of the LDOS between superconducting and normal areas, one should go beyond the mean field Usadel equation: mesoscopic fluctuations could help explain the inhomogeneous superconducting state, the re-entrant field tuned transition in TiN and modify the proximity effect.

We are grateful to M. Houzet for helping us in resolving self-consistently Usadel equation and to M. Sanquer who initiated these experiments.

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