Detection of multiple fluorescent labels using superconducting tunnel junction detectors

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(Received 27 January 2003; accepted 4 June 2003)

We show that cryogenically cooled, photon counting superconducting tunnel junctions (STJs) can be used to simultaneously record the optical spectra from multiple biological fluorochromes. Measurements with a single-pixel tantalum STJ with a wavelength resolving power \( R = \lambda / \Delta \lambda \) of about 10 confirm the expected sensitivity advantage with respect to the photomultiplier-based detectors commonly used to record signals from microarrays and other fluorescent biological systems. © 2003 American Institute of Physics. [DOI: 10.1063/1.1599059]

I. INTRODUCTION

The simultaneous measurement of the optical emission from collocated fluorochromes is a common problem in biology. Imaging cell components demands the temporal registration of multiple fluorescent markers such as variants of green fluorescent protein (GFP). In fluorescent resonant energy transfer (FRET), changes in emission spectra indicate the physical separation between donor and acceptor dye molecules. Quantifying the hybridization of labeled nucleic acids (probes) to immobilized target molecules in microarray format or in cells \textit{in situ} ideally requires the efficient, artifact-free measurement of several fluorescent spectra simultaneously.\textsuperscript{1} Cryogenically cooled superconducting tunnel junction (STJ) detectors\textsuperscript{2–6} developed for astronomy measure the energies of individual optical photons with very low levels of internal background. The operation of a STJ is indicated schematically in Fig. 1(a). In this article, we show that STJs can be used for the sensitive, quantitative registration of multiple biological fluorochromes. Using a single near-UV excitation filter, and standard dichroic and long-pass filters, the emission spectra of common fluorochromes could be easily distinguished—from each other and from broadband substrate fluorescence. STJ detectors have the potential to revolutionize quantitative multifluorochrome imaging in biology.

Photomultiplier tubes (PMTs) and image intensifiers are the photon-counting detectors most commonly used to image weakly fluorescent biological specimens. These detectors provide information on photon color (energy or wavelength) only when used with narrow band output filters or dispersive gratings. Their efficiency is limited by the quantum yield \( Q \) electrons/photon of the PMT photocathode. For a photocathode such as S20 (CsKNaSb),\textsuperscript{7} \( Q < 20\% \) in the optical band. Grating-based systems offer up to 32 parallel energy channels,\textsuperscript{8} each readout by an independent PMT, but the instrument sensitivity remains limited by the photocathode’s thermionic dark noise count rate \(- B \approx 10 \text{ counts cm}^{-2} \text{ s}^{-1} \) at room temperature.\textsuperscript{9} More commonly, the fluorochromes are addressed sequentially, repeatedly scanning the sample and registering the fluorescence through a series of narrow band output filters. Multiple scanning increases the possibility of photobleaching.

The low throughput of dispersive spectrometers was one of the motivations for the development in astronomy in the 1990’s of cryogenic, energy-resolving detectors such as the STJ and the transition edge sensor (TES).\textsuperscript{10} A STJ consists of two superconducting layers held at a temperature \( T \) well below their superconducting-to-normal transition temperature \( T_c \), and separated by an insulating aluminum oxide layer [see upper inset to Fig. 1(a)]. For \( T \sim T/10, \) the thermally induced tunnel current is negligible and the absorption of a photon of wavelength \( \lambda \) (nm) gives rise to a number \( N_0 \sim 7 \times 10^7/\lambda \Delta (T) \) of excess free charges (quasiparticles) where \( \Delta \) (meV) is the superconducting energy gap.\textsuperscript{4} The intrinsic (or “tunnel limited”) resolving power \( R = \lambda / \Delta \lambda \) of a symmetrical STJ is then

\[
R = 357(\lambda \Delta)^{-1/2} [1 + F + 1/(n)]^{-1/2};
\]

where \( F \) is the Fano factor (\(-0.22 \) for transition metal
deposited on a polished sapphire substrate. The low leakage using photolithographic techniques from a Ta/Al multilayer and the energy gap barrier. The transition temperature for bulk tantalum is 4.5 K layers and 30 nm thick Al layers on either side of the tunnel barrier. The equation for each quasiparticle tunnels across the barrier.

\[ \langle n \rangle = \frac{3}{40000} \text{tunnled electrons per eV of photon energy, and a pulse decay time of } \sim 20 \, \mu s. \] Equation (1) then indicates resolving powers of 16.0 and 8.8 at wavelengths of 600 and 2000 nm. Calibration using monochromatic radiation indicated \( R \) (600 nm) \( \sim 13.3 \). Cooling to 300 mK in a \(^3\)He cryostat (i.e., \( T \sim T_c/15 \)) kept the thermally excited quasiparticle current well below the leakage current level. We would then expect the dominant background superconductors\(^6\) and \( \langle n \rangle \) is the average number of times that each quasiparticle tunnels across the barrier.

II. MEASUREMENTS

The detector used in our initial study of biological fluorescence was a single 30×30 \( \mu m^2 \) STJ with 100 nm thick Ta layers and 30 nm thick Al layers on either side of the tunnel barrier. The transition temperature for bulk tantalum is 4.5 K and the energy gap \( \Delta(0) \) is 0.69 meV. The detector was made using photolithographic techniques from a Ta/Al multilayer deposited on a polished sapphire substrate. The low leakage current \((<0.1 \, pA/\mu m^2)\) gave a negligible contribution to the baseline noise floor, while the high transmission of the barrier resulted in high signal amplitudes. The detector had a responsivity \( \langle n \rangle N_0 \sim 40000 \) tunneled electrons per eV of photon energy, and a pulse decay time of \( \sim 20 \, \mu s. \) Equation (1) then indicates resolving powers of 16.0 and 8.8 at wavelengths of 600 and 2000 nm. Calibration using monochromatic radiation indicated \( R \) (600 nm) \( \sim 13.3 \). Cooling to 300 mK in a \(^3\)He cryostat (i.e., \( T \sim T_c/15 \)) kept the thermally excited quasiparticle current well below the leakage current level. We would then expect the dominant background

source in the STJ to be cosmic ray muons interacting with the sapphire substrate, in a complex manner, to create pair-breaking phonons at an upper limit rate \( B \sim 0.01 \, cm^{-2} \, s^{-1}. \) Given a peak STJ quantum efficiency \( Q \) (600 nm) of 75%, we then arrive at the sensitivity advantage of the cryogenic detector over a PMT-based system:

\[
\frac{Q_{\text{STJ}}/\sqrt{R_{\text{STJ}}}}{Q_{\text{PMT}}/\sqrt{R_{\text{PMT}}}} = 120 \text{ times.}
\]

This figure could ultimately be higher in a many-pixel STJ array, where substrate events would give rise to simultaneous pulses in adjacent pixels.

Light from a Leica Aristoplan epifluorescence microscope with a 100 W Ushio HBO mercury excitation source was coupled into the backilluminated STJ via a 6 m Oriel 77530 UV-grade fused silica fiber, bandpass \( \lambda \sim 200\text{–}2000 \) nm. The STJ quantum efficiency exceeds 50% for \( \lambda \) values between 150 and 700 nm. No efforts were made to focus the light from the specimen onto the fiber or from the end of the fiber onto the detector. The coupling efficiency was therefore extremely low \((<0.01\%).\) The detector was held within a light tight shield at the base system temperature of 300 mK. The warm end of the optical fiber, however, constitutes a 300 K black body source which, when convolved with the fiber transmission, provides an effective reference signal in all STJ spectra at a wavelength of 2 \( \mu m \) (see Figs. 2 and 4).

The warm readout electronics consisted of a charge sensitive preamplifier situated \( \sim 1 \, m \) from the detector, a shaping amplifier (10 \( \mu s \) shaping time) and an analog to digital (AD) converter whose output was optically coupled to a PC, on which the pulse amplitude and decay time was recorded for each detected photon. A pulse risetime window was applied to select valid events.

Fluorescence spectra were measured from eleven different, commercially available, labeled uridine nucleotides. Six different substrates were also examined. Four excitation filters were used: the spectra presented here were obtained with the following two filter sets:

(i) Leica filter set I3; 450–490 nm excitation, 510 nm dichroic cutoff, and 515 nm long-pass filter.

(ii) Omega Optical Inc.\(^{11}\) 56 triple excitation filter, with transmission bands centered on 457, 528, and 633 nm, its associated dichroic mirror and an Omega 580 nm long-pass emission filter.

The transmissions of the actual filters used were confirmed \textit{post facto} using a laboratory spectrophotometer.

III. ANALYSIS

Even though excited far from maximum absorbance, spectra were obtained from fluorochromes emitting in the red (Alexa 594, Fluorored, and Texas Red) and green (Fluorogreen, Avidin FITC, and Alexa 488). We illustrate the potential of the STJ for fluorescent imaging with particular reference to the dyes Alexa 488 and Alexa 594.\(^{12}\) Figure 2 shows the pulse height spectrum of the latter fluorochrome with the Leica I3 filter set. The spectrum is cutoff at 2.4 eV (515 nm) by the long-pass filter. Even with the fluorochrome excited at only \( \sim 3\% \) of maximum absorbance, the signal-to-
background (peak channel count divided by average channel count at energies just above the IR peak) is at least 275:1. Fitting a normal distribution (the broken curve) to the IR peak indicates $R (2000 \text{ nm}) = 4.0$. Injecting a known charge to the warm STJ preamplifier from an electronic pulser produced a peak width of 0.084 eV FWHM. The quadrature sum of tunnel limit and electronic noise contributions then implies a 2 $\mu$m resolving power $R = 5.7$, slightly better than observed.

For each fluorochrome, the microscope focus was adjusted to produce an output count rate of 800 Hz—the maximum rate before pulse pileup compromised the spectral resolution. The “dark” count rate obtained by closing the microscope shutter between bulb and filter was 0.09 s$^{-1}$. Because individual spectra were not obtained at constant intensity, overlaying two STJ spectra does not necessarily represent the degree of spectral separation for the corresponding mixture of fluorochromes. Nevertheless, Fig. 3 indicates that, while the degree of spectral overlap is significant, it is straightforward to deconvolve the relative intensity contributions of mixed Alexa 488 and Alexa 594 probes. We note that the peak emission energies and spectral widths for both probes differ significantly from the templates published by the manufacturer.$^{12}$ The peak emission energy for Alexa 594 is redshifted by 0.17 eV from the expected value of 2.088 eV; that of Alexa 488, by 0.29 eV. The measured width of the Alexa 594 distribution is somewhat larger than obtained from the convolution of the manufacturer’s spectrum with the energy response of the STJ. Since the STJ calibration rules out any “zero shift” in the detector response (see Fig. 4), we conclude that these measurements constitute evidence for the influence of local environment on fluorochrome emission.
sion. Here, the dyes dried out from solution spotted onto glass cleaned with chromic acid.

Figure 5 shows the Alexa 594 and Alexa 488 spectra obtained with the Omega triple filter set; the “top hat” transmission bands of the output filter are indicated by the broken vertical lines. This figure shows that the spectroscopic capability of the STJ may be combined with appropriate selections of input and output filters to give highly preferential registration of one fluorochrome. In band 1, centred on a photon energy of 2.5 eV, the ratio of peak intensities is about 10:1, while in band 2 at 2.8 eV, the ratio is 1:1.

Figure 6 illustrates the first application of the STJ to the study of genetic material hybridized, on an Arrayit™ substrate, with multiple probes (Alexa 568, Alexa 594, and Bodipy 630/650—all produced by Molecular Probes Inc.). Data set 1 represents the measured sample spectrum, while curves 2, 4, 5, and 6 are the component spectra measured for the three fluorochromes separately and for the substrate. Curve 3 indicates that a linear sum of the component intensities with weights:

\[
\text{Alexa 568}:\text{Bodipy 630/650}:\text{Substrate}:\text{Alexa 594} = 10:10:4:3,
\]

provides an excellent fit to the data in the 1.9–2.4 eV (515–652 nm) band. We cannot attribute the excess intensity at lower energies to the presence in the sample of unhybridized DNA, since the intrinsic fluorescence of DNA is well known to peak in the UV. One explanation is excimer (excited dimer) emission arising as a result of the high concentrations of the fluorochrome probes in the sample, although fluorescence from components of the solutions used to prepare the array may also be contributing. The manufacturer’s handbook for Bodipy dyes states that these fatty acid analogs do generally exhibit concentration-dependent excimer formation and consequent redshifting of their fluorescence.

IV. DISCUSSION

We have shown that a cryogenic detector can detect, quantitatively and at very high signal-to-noise compared to...
conventional systems, characteristic fluorescent emission spectra from fluorochromes commonly used in biology. Even with single excitation wavelength bands, photon-by-photon registration of the complete spectrum allows separation of multiple fluorochromes. Remarkably, we found that nonfocussed, low levels of peripheral sample illumination with Philips TLD 36W/83 fluorescent tubes (i.e., normal laboratory lighting) were able to excite count rates in excess of 1000 Hz from fluorochromes on a slide on a microscope stage.

The present Ta technology ($R \sim 10^{-20}$) is compatible with at least four simultaneous labels with a single excitation filter; the potential of Hf [$\Delta(0) = 0.02$ meV; $R \sim 80$] and Mo [$\Delta(0) = 0.14$ meV; $R \sim 40$] for better resolution (at lower operating temperatures) is well documented.\(^4\) In the context of biological assays, in particular the measurement of hybridization of labeled probes to microarrays, any increase in the number of labels that can be measured without interference is important because either increased numbers of internal controls can be included, or more samples can be tested simultaneously. The modest count rate limitation of a single STJ pixel can be overcome by using parallel arrays; 6×6 element Ta STJ matrices have been fabricated and the development of 10×32 Mo arrays studied by the ESTEC Group [see Fig. 1(b)]. These developments and the production of closed cycle cooling systems for the $T<100$ mK regime will have an important bearing on the ultimate utility of optical STJs in the life sciences.

**ACKNOWLEDGMENT**

The authors acknowledge the support of Muna Naji in reducing the STJ datasets.

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1. T. Schwarzacher and J. S. Heslop-Harrison, *Practical In Situ Hybridization* (Bios, Oxford, 2000), p. 213.
2. M. A. C. Perryman, C. L. Foden, and A. Peacock, Nucl. Instrum. Methods Phys. Res. A 325, 319 (1993).
3. A. Peacock, P. Verhoeve, N. Rando, A. van Dordrecht, B. G. Taylor, C. Erd, M. A. C. Perryman, R. Venn, J. Howlett, D. J. Goldie, J. Lumley, and M. Wallis, Nature (London) 381, 135 (1996).
4. P. Verhoeve, N. Rando, A. Peacock, A. van Dordrecht, A. Poelaert, and D. Goldie, IEEE Trans. Appl. Supercond. 7, 3359 (1997).
5. N. Rando et al., Rev. Sci. Instrum. 71, 4582 (2000).
6. N. Rando, A. Peacock, A. van Dordrecht, C. L. Foden, R. Engelhardt, B. G. Taylor, J. Lumley, and C. Pereira, Nucl. Instrum. Methods Phys. Res. A 313, 173 (1992).
7. A. H. Sommer, *Photoemissive Materials* (Wiley, New York, 1968).
8. META detector for the Carl Zeiss LSM 510 Confocal Microscope. Carl Zeiss, Welwyn Garden City, Herts, UK.
9. I. G. Butler et al., Proc. SPIE 2278, 126 (1994).
10. R. W. Romani, A. J. Miller, B. Cabrera, S. W. Nam, and J. M. Martinis, Ap. J. (in press).
11. Omega Optical Inc., 210 Main St., Brattleboro, VT 05301.
12. N. Panchuk-Voloshina et al., J. Histochem. Cytochem. 47, 1179 (1999); Molecular Probes Inc., P.O. Box 22010, Eugene, OR 97402-0469.
13. TeleChem International Inc., 524 E. Weddell Dr., Suite No. 3, Sunnyvale, CA 94089.