Low kinetic inductance superconducting MgB$_2$ nanowires with a 130 ps relaxation time for single-photon detection applications

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Received 29 October 2020, revised 21 December 2020
Accepted for publication 22 January 2021
Published 23 February 2021

Abstract

Properties of superconducting nanowires set the performance level for superconducting nanowire single photon detectors (SNSPDs). Reset time in commonly employed large area SNSPDs, 1–10 ns, is known to be limited by the nanowire’s kinetic inductance to the load impedance ratio. On the other hand, reduction of the kinetic inductance in small area (waveguide integrated) SNSPDs prevents biasing them close to the critical current due to latching into a permanent resistive state. In order to reduce the reset time in SNSPDs, superconducting nanowires with both low kinetic inductance and fast electron energy relaxation are required. In this paper, we report on a study of kinetic inductance in narrow (15–100 nm) and long (up to 120 $\mu$m) superconducting MgB$_2$ nanowires made from 5 nm thick films, offering such combination of properties. Such films were grown using hybrid physical chemical vapor deposition, resulting in a critical temperature of $\sim$32 K, and a switch current density of $5 \times 10^7$ A cm$^{-2}$ (at 4.8 K). Using microwave reflectometry, we measured a kinetic inductance of $L_{k0}(4.8$ K) = 1.3–1.6 pH/□ regardless of the nanowire width, which results in a magnetic field penetration depth of $\sim$90 nm. These values are very close to those in pristine MgB$_2$. We showed that after excitations by a 50 fs pulsed laser the reset time in 35 nm $\times$ 120 $\mu$m MgB$_2$ nanowires is 130 ps, which is more than a factor of 10 shorter than in NbN nanowires of similar length-to-width ratios. Depending on the bias current, such MgB$_2$ nanowires function as single-, double, or triple-photon detectors for both visible ($\lambda = 630$ nm) and infrared ($\lambda = 1550$ nm) photons, with a dark count rate of <10 cps. Although the apparent photon detection efficiency seems so far to be low, further technological advances (uniform nanowire width, smaller thickness, increasing the switching current closer to the pair-breaking current) may improve this figure of merit.

Keywords: MgB$_2$, kinetic inductance, superconducting nanowire, SNSPD, thin superconducting film, single photon detector

(Some figures may appear in colour only in the online journal)
1. Introduction

During the last two decades we have seen enormous progress with superconducting nanowire single photon detectors (SNSPDs), from first demonstration of principles [1] to integrated circuits and a wide range of applications in nanophysics [2], high speed communication [3], quantum information [4], quantum key distribution [5], laser-ranging [6], deep space communication [7], etc. SNSPD’s operation is based on photon assisted disruption of supercurrent in a superconducting nanowire, which is dc-biased close to the critical current [8, 9]. Because the superconducting energy gap Δ is in the meV-range, a large number of non-equilibrium quasiparticles are generated at visible or infrared photon (~eV) impacts. Both the small width of the nanowire, w, and the critical current, Ic, being close to the pair-breaking current ensure high detection efficiency [10, 11]. After hot-spot formation, the bias current is diverted from the SNSPD into the readout. Within a relaxation time, τ0 (set by electron-phonon non-elastic scattering and phonon escape into the substrate), non-equilibrium quasiparticles recombine into Cooper pairs. Until the supercurrent in the nanowire is restored and the hot-spot disappears (the reset time), SNSPD is blind to any further incoming photons. NbN SNSPDs, being the first one where single-photon sensitivity was demonstrated, still provide the shortest reset time for a given detector area, compared to SNSPDs made from other superconductors. In 5 nm thick NbN films, the quasiparticle relaxation time τ0 is ~30–50 ps [12, 13]. However, in long (l) and narrow (w) NbN nanowires, large kinetic inductivity (Lk, 4.8 K = 80–90 pH) (inductance per square of the nanowire, l = w) leads to a significantly slower current return (restoration of the superconducting state), resulting in a much larger reset time τ = Lk/Rk ≫ τ0, where Lk = Lk0 × l/w is the total kinetic inductance of the nanowire and Rk is the impedance of the read-out circuit (~50 Ω). This is particularly critical in large area SNSPDs, with a kinetic inductance as large as hundreds of nH, leading to a reset time of a few tens of ns. On the other hand, large kinetic inductance delays current return from the load Rk back into the SNSPD, prior to the hot-spot cooling down, hence preventing the SNSPD from latching into a permanent normal (non-sensitive) state [15]. To a certain extent, the reset time can be reduced by choosing a high-impedance load. However, much increased Rk leads to a reduction of the maximum non-latching bias currents, hence precluding SNSPDs from reaching high detection efficiencies [16]. This problem is particularly pronounced in short SNSPDs, where shunting resistors of small values or extra inductors in series are required to avoid latching, which does not altogether allow reaching short reset times [17, 18].

Apart from low-Tc superconductors (NbN, Nb, NbTiN, TaN, MoN, WSi, MoSi, etc) [10, 15, 19–21], magnesium diboride (MgB2) nanowires have also shown ability for single photon detection in both visible [22] and IR [23] ranges. However, the reset time in those MgB2 SNSPDs was reported to be 2–3 ns, i.e. close to that for NbN SNSPDs, the fact that was left unexplained. Previously it has been shown that in clean MgB2 (attainable in 50–200 nm thick films) the magnetic field penetration depth λ0 is at least an order of magnitude shorter than in NbN [24], which should lead to a small kinetic inductivity (Lk0 = μ0λ0/2Ic). Apparently, due to a large number of defects in the abovementioned thin MgB2 films (normal state resistivity ρ0 > 300 μΩ cm) kinetic inductance increases, which affects the reset time. Recently, we managed to deposit 5 nm thick MgB2 films with a critical temperature >30 K and ρ0 comparable to that in thick films. Furthermore, in such thin films, a quasiparticle relaxation time is measured to be τ0 = 12 ps [25], and which all together suggests that MgB2 nanowires might provide a solution for high speed photo detection.

In this work, we study both kinetic inductance and response rate in nanowires, with a large variation of width (15–900 nm) and length (3–120 μm), made from clean (low normal state resistivity) and ultra-thin (5 nm) MgB2 films. Observing a kinetic inductance as low as 1.5 pH, we show experimentally that a 130 ps voltage relaxation time can be achieved in 120 μm-long devices into a standard 50 Ω readout circuit with a 100 MHz repetition rate. When biased close to the critical (switch) current, such devices show a linear dependence of the number of response pulses vs the incident laser power, corresponding to the single-photon detection regime.

2. Film deposition and device fabrication

MgB2 films were grown on 6 H-SiC substrates using a custom-built hybrid physical chemical vapor deposition (HPCVD) system [26, 27]. The system was designed for a low deposition rate (~2.5–3 nm min−1) with a diborane gas flow of 2 sccm (5% B2H6 in H2) and a deposition temperature of 700 °C. As a result, continuous MgB2 films could be routinely obtained as thin as 5 nm, as shown by transmission electron microscopy (TEM) (figure 1(a)), with a critical temperature of 32–34 K (figure 2(a)). Film growth on the SiC substrate is epitaxial, which ensures that bulk properties of MgB2 are preserved in the thin films as manifested in a fairly low normal-state (residual) resistivity of ρn = 10–15 μΩ cm, a high switch current density of Jc ~ 5 × 107 A cm−2, and a small magnetic field penetration depth of λ = 90 nm (see further in the text). TEM analysis was conducted on a set of MgB2 films of varying thickness, where the thickness proportionality to both the diborane flow and the deposition time was verified.

Subsequently, MgB2 films were transferred (vacuum broken) into a dc-magnetron sputtering system where plasma assisted cleaning of the film surface was done followed by in situ sputtering of a 20 nm thick Au film. This was followed by the device fabrication route #1 which starts with defining the coplanar waveguide (CPW) contact pads and alignment marks on the MgB2 films with a 20 nm gold layer. A layer of polymethyl methacrylate (PMMA—70 nm) on top of copolymer (MMA—470 nm) was utilized as a positive tone bilayer resist system for electron beam lithography. The subsequent development in isopropanol/water mixture (10:1), electron beam physical vapor deposition of Ti–Au–Ti (10 nm–200 nm–40 nm), and lift-off in acetone leaves the desired patterns on the substrate. The first 10 nm titanium layer
promotes the adhesion of Au whereas the final 40 nm layer serves as a mask for gold contact pads later on during the Ar$^+$ ion milling. The nanowire patterns were defined via another electron beam lithography step with a negative tone resist (ma-N 2401—100 nm), developed in tetramethlyamonium hydroxide aqueous solution (ma-D 525) followed by rinse in DI water. The remaining MgB$_2$ and the initial 20 nm Au layer, except on top the nanowire, were etched using Ar$^+$ ion milling. Finally, the negative resist mask layer was removed with acetone and the initial 20 nm Au layer on top of the nanowire was removed by another Ar$^+$ ion-milling step (figures 1(b) and (c)).

A second fabrication route has also been explored which differs from the route #1 by absence of the initial gold protection layer (20 nm Au in Route #1) on top of the MgB$_2$ film. In this case, the final ion milling step, as in the case of route #1, is unnecessary, therefore there is no need in removing the negative resist mask and hence Au layer. The route #2 contains fewer steps, whereas we have not observed any MgB$_2$ film degradation due to absence of the protective Au layer.

In order to compare both the kinetic inductance and the reset time to those in a well-established material system, we have also fabricated and tested 100 nm wide NbN nanowires, both 115 µm and 460 µm long. The NbN film (d ∼ 5 nm thick, as estimated from the deposition rate) was deposited by DC reactive magnetron sputtering in an Ar + N$_2$ environment on a 330 µm thick C-plane double side polished sapphire substrate maintained at 800 °C. The film was patterned into 100 nm wide (100 nm spacing) meandering nanowires with reactive ion etching (CF$_4$:O$_2$, 5:1 flow ratio) through a mask defined by e-beam lithography on a positive (PMMA—80 nm) resist. During the RIE process, the total pressure was kept as low as possible (3.8 mTorr being the lowest used to provide a stable plasma discharge) in order to increase the etching anisotropy. Electrical contact pads (as single-port 50 Ω CPW) were fabricated in 10 nm Ti/200 nm Au/40 nm Ti using a standard lift-off process. The NbN film between the CPW contacts was removed by means of wet-etching (nitric acid/hydrofluoric acid/acetic acid, 5:3:3) with a resistive mask placed over the nanowires (photo resist AZ1512). The critical temperature was ∼10 K and the room temperature sheet resistance was ∼300 Ω/□, which corresponded fairly well to NbN film characteristics for previously published NbN SNSPDs.

### 3. Experimental Techniques

All experiments were conducted in a cryogen-free probe station Lakeshore CRX-4K (base temperature 4.8 K) with an optical view port and microwave ground-signal-ground probes on XYZ manipulator stages (figure 3(a)). Kinetic inductance was obtained from the imaginary part of

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**Figure 1.** (a) TEM image of a 5 nm-thick MgB$_2$ film on the SiC substrate. (b), (c) SEM images of a meandering MgB$_2$ nanowires. Scale bars are given in each of the images.

**Figure 2.** (a) Superconducting transitions (normalized resistance) in an MgB$_2$ and an NbN nanowire. (b) I(V)-curves (at 5 K) for a MgB$_2$ and a NbN nanowire, utilized for the response rate experiments (see figure 5). Bias points are indicated with arrows. The inset shows simplified schematics of a photon detector (a variable resistance and an inductor, representing kinetic inductance in the nanowire) with a biasing network and a readout load (e.g. a low-noise amplifier).
the microwave impedance \( Z(f) = 2\pi f L_k \) where \( f \) is the probing microwave frequency), which was measured using a (10 MHz–67 GHz) Vector Network Analyzer Rohde&Schwarz ZVA 67. The vector network analyzer (VNA) calibration was done using a wafer with calibration standards (matched (50 Ω), open, and short) placed on the same cooling stage next to the MgB\(_2\) samples. Therefore, both the VNA calibration and impedance measurements of MgB\(_2\) nanowires were performed in a single cool-down. Magnetic (or geometric) inductance in superconducting films as thin as those utilized in our experiments is negligibly low, and it can be calculated analytically using the approach from [28] as a function of the film thickness \( d \) and the temperature \( T \):

\[
L_m(d, T) = \frac{\mu_0 \lambda_0(T)}{4} \left[ \coth \left( \frac{d}{2\lambda_0(T)} \right) - \frac{d}{2\lambda_0(T)} \right] - \text{csch} \left( \frac{d}{2\lambda_0(T)} \right)^2
\]

where \( \mu_0 \) is the vacuum permeability and \( \lambda_0(T) \) is the magnetic field penetration depth. Considering the \( d = 5 \) nm and \( \lambda_0(5 \text{ K}) = 90 \) nm, \( L_m(5 \text{ nm, } 5 \text{ K}) = 0.5 \) pH/□.

Lasers flood-illuminated the samples through a quartz pressure window of the probe station. The standard IR filter (not transparent for 1560 nm photons) on the 60 K shield was replaced with a WG12012-C window (Thorlabs). The output beam from the cw \( \lambda = 630 \) nm laser diode module required no extra collimation, resulting in a light spot diameter of \( \sim 4 \) mm. The IR (\( \lambda = 1560 \) nm, Toptica FemtoFerb 1560, 100 MHz repetition rate) 50 fs pulse laser is fiber-coupled and a fiber launcher was utilized to collimate the IR laser beam. The IR laser spot size (on the sample) was approximately 5 mm, verified using SNSPD photo counts vs the laser XY-shift. Two sets of light attenuators were utilized for the visible and IR light. The emission of both lasers (\( \sim 2 \) mW and \( \sim 100 \) mW (average) for the \( \lambda = 630 \) nm and \( \lambda = 1560 \) nm lasers, respectively) was attenuated to a level where the switch current in the nanowires is not visible anymore.

At room temperature, both microwave readout and dc biasing were arranged (figure 3(b)) though a bias-T (Picossecond 5547, 12 kHz–15 GHz) followed by a low noise amplifier (MITEQ 3D-0010025, 0.5–5 GHz) and either a real time oscilloscope (Keysight Infinium 54854 DSO, 4 GHz, 20 GSa s\(^{-1}\)) or a pulse counter (HP 53131A 225). Nanowires were biased with a Yokogawa 7651 programmable dc source.

4. Kinetic inductance

As measured in a large variety of samples (width, length and fabrication batches), the kinetic inductivity in thin MgB\(_2\) nanowires was 1.35–1.60 pH/□ at 4.8 K (figure 4(a)). In the case of defect-free uniform films, the total kinetic inductance \( L_k = L_k^{\text{F}} + L_k^{\text{G}} \) is expected to be proportional to the length-to-width ratio \((l/w)\) and hence to the total resistance \( R \) in the normal state (40 K). \( R_s \) is the MgB\(_2\) film sheet resistance in the normal state. This trend was indeed observed (figure 4(b)), which confirms scalability of thin-film transport properties from the micron- to the nano-scales.

Kinetic inductance (and its temperature dependence) is a sensitive indicator for superfluid (Cooper pairs) density, hence providing the value for the superconducting energy gap. In MgB\(_2\), \( \pi \)- and \( \sigma \)-sheets of the Fermi surface display quite dissimilar energy gaps \( \Delta_\pi(T) \) and \( \Delta_\sigma(T) \), with temperature dependences following [29]:

\[
\Delta_\pi(T) = \Delta_\pi(0) \left[ 1 - \left( \frac{T}{T_c} \right)^{1.8} \right]^{1/2}
\]

\[
\Delta_\sigma(T) = \Delta_\sigma(0) \cdot \left[ 1 - \left( \frac{T}{T_c} \right)^{2.9} \right]^{1/2}
\]

\( \Delta_\pi(0) \) and \( \Delta_\sigma(0) \) are zero temperature energy gaps in \( \pi \)- and \( \sigma \)-sheets.

Previously, in the frame of double-band modeling [30], it has been shown that electrodynamic properties in MgB\(_2\) can be analyzed by considering two quasi-layers (electrons of both \( \pi \)- and \( \sigma \)-bands) in parallel, with a finite (yet, weak) inter-band scattering. In particular, partial contribution from each band to both the condensate energy and the superfluid density varies according to temperature and magnetic field, e.g. at zero magnetic field, the \( \pi \)-band is expected to contribute about \( a > 70\% \) to the superfluid density (\( \times \sqrt{\lambda_0\Delta_0} \)) at \( T < (T_c - 2 \text{ K}) \).

The exact value of the coefficient \( a \) depends on the interband scattering rate, and has to be evaluated for each particular case. However, the general trend seems to be correct, as in [24], where it was shown that \( \lambda_0 \) vs the mean free path can be modelled considering predominantly the \( \pi \)-band. In [31], a satisfactory agreement with experimental \( \lambda_0(T) \) was demonstrated for thick films, considering two bands with an assumed \( \pi \)-band weighing coefficient \( a = 0.8 \). For a general case of two conduction bands, the kinetic inductivity can be calculated from the imaginary part of the BCS conductivity [32] as:

\[
L_{k0}(T) = \left[ \frac{\hbar \cdot 2R_s}{a \cdot \pi \cdot \Delta_\pi(T)} \cdot \frac{1}{\tanh \left( \frac{\Delta_\pi(T)}{2k_B T} \right)} \right]^{-1} + \left[ \frac{\hbar \cdot 2R_s}{(1-a) \cdot \pi \cdot \Delta_\sigma(T)} \cdot \frac{1}{\tanh \left( \frac{\Delta_\sigma(T)}{2k_B T} \right)} \right]^{-1}
\]

\( k_B \) is the Boltzmann constant, and \( \hbar \) is the 2\( \pi \)-normalized Planck constant.

By fitting equation (2) to the experimental \( L_{k0}(T) \) data (figure 4(c)), we obtained the superconductor energy gaps at \( T = 0 \). Our results showed that equation (1) holds even for extremely thin MgB\(_2\) films considering that \( \Delta_\pi(0) \) is 0.60\( k_B \cdot T_c = 1.7 \) meV and \( \Delta_\sigma(0) \) is 2.22\( k_B \cdot T_c = 6.2 \) meV, with an interband coefficient \( a = 0.85 \). For a mean experimental value of \( L_{k0}(5 \text{ K}) = 1.5 \) pH/□, we can obtain the effective magnetic field penetration depth: \( \lambda_0 \)
of the nanowire response were 90 ps and 130 ps (10%–90%),
100 MHz repetition rate of the laser. The rise and the fall time
readout. The measured voltage impulses corresponded to the

demonstrate a reset time (10%–90%), which is at least a factor
10 faster than in NbN nanowires. As follows from figures
(a) and (b), MgB$_2$ nanowires demonstrate a reset time (12 ps in
5 K) and (a), both corresponding well to the measured kinetic

depth. Small kinetic inductance in MgB$_2$ nanowires is a very
attractive feature in SNSPD applications. For one of the
samples, corresponding to an aver-

tation depth for the measured resistivity of
(5 K) [34]. Increase of the penetration
depth in the clean limit [32]. The BCS coherence length can be estimated from
$\xi_{\Omega} = h\nu_e/(\pi \Delta_\pi) = 66 \text{ nm}$, considering the $a$–$b$ plane averaged Fermi velocity as $\nu_F \sim 5.4 \times 10^5 \text{ m s}^{-1}$ [30]. For ultrathin films, as utilized in our study, electron scattering on the film surface might become significant (i.e. $\ell \sim 5 \text{ nm}$), hence increasing both the residual resistivity and the penetration depth.

Small kinetic inductance in MgB$_2$ nanowires is a very
attractive feature in SNSPD applications. For one of the
longest (in the number of squares, i.e. the $L/w$ ratio) of the
utilized in our study MgB$_2$ samples (35 nm × 120 μm, 3428
squares) we measured its impulse response on excitations from
a 50 fs pulsed laser ($\lambda = 1550 \text{ nm}$). The measured total inductance for this sample was 5100 pH, which is expected to result in a voltage fall time of $\tau = L_d/R_l = 102 \text{ ps}$ for a 50 Ω readout. The measured voltage impulses corresponded to the 100 MHz repetition rate of the laser. The rise and the fall time of the nanowire response were 90 ps and 130 ps (10%–90%), respectively (figures 5(a) and (b)). The observed fall time is close to that calculated from the kinetic inductance.

For comparison, both the kinetic inductance and the
impulse response were measured for two types of 100 nm wide NbN nanowires, 115 μm (1150 squares) and 460 μm (4600 squares) long. The kinetic inductance was measured in these samples (54 nH and 200 nH, respectively) using the same method as for MgB$_2$ samples, corresponding to an average $L_{\Omega}(5 \text{ K}) = 45 \text{ pH}$. This value is $\sim 1/2$ of the previously reported [14], which can be due to variations of the NbN film quality (choice of the substrate, deposition conditions, etc), due to uncertainty in the film thickness, as well as due to variations of $L_\pi$ measurements techniques. The voltage impulse decay time in NbN nanowires was 4.1 ns for the 460 μm long sample and 1.3 ns for the 115 μm long sample (figures 5(a) and (b)), both corresponding well to the measured kinetic inductance.

As follows from figures 5(a) and (b), MgB$_2$ nanowires demonstrate a reset time (10%–90%), which is at least a factor of 10 faster than in NbN nanowires of a similar geometry. Despite of being biased at the switch current, MgB$_2$ nanowires with a kinetic inductance of 5 nH demonstrate no latching, which follows from the fast decaying voltage impulses (figures 5(a) and (b)). Fast quasiparticles relaxation (12 ps in MgB$_2$ vs 50 ps in NbN) [25] is a positive factor which reduces the effect of latching in small kinetic inductance samples.
The voltage impulse rise dynamics in superconducting nanowire detectors has previously been studied both experimentally and theoretically in SNSPDs made from NbN [14, 33, 34]. One of the conclusions made there was that the rise time is proportional to the square root ratio of the total inductance and the normal domain resistance \( L_{\text{tot}} \sim \sqrt{R_{\text{ND}} + l_0} \), approaching nearly linear dependence at high inductance values \( L_{\text{tot}} \sim \frac{l_0}{R_{\text{ND}} + l_0} \). Because the sheet resistance in NbN is of the order of 500 \( \Omega \), the rise time in NbN SNSPDs is much less than the fall time. The sheet resistance in thin MgB\(_2\) films is 20–30 \( \Omega \), which can lead to a rise time longer than in NbN for the same total inductance. Considering that in MgB\(_2\) the kinetic inductance is a factor of 30–60 lower than in NbN, the increased rise time will be noticed only for very long samples. Detailed modeling and experimental studies are needed in order to clear the question of hot-spot formation in MgB\(_2\) nanowires and to understand what will be the limiting factor for the rise time both for short and long samples.

For an initial estimate of the MgB\(_2\) photon detector response jitter, we utilized statistics for response pulse intervals collected within a 10 \( \mu \)s time-slot through a single oscilloscope scan, which was triggered by the first detection event. The response interval jitter is still \~50 ps, which corresponds to a \~25 ps jitter of the rising edge (figure 5(c)). This value is an upper limit for MgB\(_2\) nanowire detectors, because a jitter less than that in NbN could be expected in MgB\(_2\) considering a factor of 33 lower kinetic inductance [33].

5. IR- and visible light detection

Direct demonstration of single photon detection capability requires a 100% photon coupling and a known quantum efficiency. Considering that the former is a challenging engineering task and the latter would require complex modelling, the single photon detection mode is often verified by statistical analysis of the pulse counts per photon flux [11] or inter-counts intervals [11]. We chose the first method, utilizing initially a cw laser (630 nm). The photon count rate \( N \) vs the laser attenuation \( F \) is shown in figure 6(a) at several bias currents for a 90 nm wide and 40 \( \mu \)m long nanowire. For bias currents \( I_0 \gtrsim 396 \mu \)A, the pulse count is linearly proportional to the photon flux, \( N \propto F^m \). For smaller bias currents we observed that \( N \propto F^m \), where \( m > 1 \), which indicates that multiphoton detection mode becomes dominant. These results show that, in contrast to NbN nanowire detectors, the single photon detection mode is limited in MgB\(_2\) nanowires to a narrow bias current range in the vicinity of \( I_c \) (figure 6(b)). Physical modeling using accurate material parameters is required in order to obtain a full understanding of photon detection in MgB\(_2\) thin film nanowires. In this moment, we note that the aforementioned fact is probably caused by a smaller size of the normal domain \( R_{\text{ND}} \) (compared to the nanowire width), formed by the photon-deposited energy. Its order of magnitude could be estimated using an expression \( R_{\text{ND}} \approx \frac{\mu \text{m}}{4\pi d N(0) (h k T)^2} \) for a photon energy \( E_{\text{ph}} \) [9]. The total density of electronic
states $N(0)$ at the Fermi level in MgB$_2$ can be obtained e.g. from the electron specific heat data as $\frac{\gamma_n}{\rho_m} = 2.3 \times 10^{47} \text{m}^{-3} \text{J}^{-1}$, where $\gamma_n = 2.6 \text{mJ mol}^{-1} \text{K}^{-2}$ is the electron specific heat coefficient in the normal state [29], $\rho_m = 2.6 \text{g cm}^{-3}$ is the mass density, and $M = 45.9 \text{g mol}^{-1}$ is the molar mass. The obtained $N(0)$ is close to that in NbN [35], whereas $T_c$ in MgB$_2$ is a factor of 3 higher. Therefore, a higher dc current ($\sim I_c$) is required to extend the normal domain to the edges of the nanowire. The fact that we have not observed any rise of the kinetic inductance at bias current close to the switch current most probably indicates that the switching current in our MgB$_2$ nanowires is conduction-limited [36]. The observed dark count rate in the studied MgB$_2$ nanowire detectors was very small, <10 cps at the highest utilized bias current (398 $\mu$A), despite the fact that the samples were not shielded from either the electrical or magnetic field interference. At such bias current the dark count rates in NbN SNSPDs often reach values of $>10^3$–$10^5$ cps [34, 35]. It is another indication that the switch current in our samples is limited by a constriction in a short section on the nanowire, leaving the most of the nanowire under-biased.

Similar results were obtained for detection of IR photons ($\lambda = 1560$ nm) from the 50 fs pulsed laser by a 35 nm $\times$ 120 $\mu$m MgB$_2$ sample ($I_e = 97$ $\mu$A). For $I_0 > 0.97 \times I_e$, a linear $N(F)$ dependence (figure 6(c)) indicates that the detector response is triggered by single photon absorption events. For $I_0 < 0.97 \times I_e$, a combination of both one- and two-photon responses was observed ($N \propto F^m$, $m > 1$). At a high photon flux ($F > 0.1$), i.e. at a large number of photons per pulse, multi-photon response even dominates at the highest bias current, as has been observed in NbN SNSPDs [37] for large laser power.

6. Conclusions

In our study, we investigated ultra-thin MgB$_2$ films in an attempt to create a new material basis for SNSPDs. Previously published works have demonstrated that single photon sensitivity is achievable with MgB$_2$ nanowires. However, the reported reset times were >1 ns even for relatively short devices, which we interpret to result from a high degree of disorder in those films, manifested in both high normal state resistivity and high kinetic inductance. HPCVD was known to be a way of making clean MgB$_2$ films, which so far been limited to thicknesses >10–15 nm. MgB$_2$ films thinner than that were reported to break into a set of isolated islands (Volmer–Weber growth mode). Using our custom-built HPCVD system we have previously observed that by reducing the deposition rate to its lowest limit ($2.5–3$ nm min$^{-1}$) MgB$_2$ films as thin as 5 nm can be deposited, which allowed to fabricate 1 $\mu$m$^2$ devices (hot-electron bolometer, HEB, terahertz mixers) with a high switch current density. In the current paper, we show that even nanowires as narrow as 20 nm and with a large aspect ratio ($L/I_w > 3000$) can be fabricated using e-beam lithography and Ar$^+$-ion milling using such MgB$_2$ films. The obtained kinetic inductance is $\sim L_0/(4.8 \text{K}) = 1.3–1.6 \text{pH} / \text{cm}$ corresponding to a magnetic field penetration depth of $\lambda_0 = 90$ nm. Scalability of both the kinetic inductance and the normal-state resistivity from micrometer down to nanometer device scale indicates a high film uniformity. High quality of films is also manifested in a high critical temperature ($>30$ K), and a high switch current density ($\sim 5 \times 10^7$ A cm$^{-2}$), approaching some of the highest values ever achieved in MgB$_2$ films. With the reset time being limited by kinetic inductance, MgB$_2$ nanowire detectors are expected to operate a factor of $L_0/(\text{NbN})/L_0/(\text{MgB}_2) = 30–60$ times faster compared to NbN devices. We confirm it experimentally by observing a voltage impulse decay time of 130 ps for an MgB$_2$ nanowire with a total kinetic inductance of 5 nH, whereas this figure is $\sim 4$ ns for an NbN nanowire with a similar $L/I_w$ ratio. Obtained results show that MgB$_2$ nanowires are capable of single photon detection in both visible and communication ($1.5 \mu$m) spectral ranges, combined with a low dark count rate, hence forming a technological platform for sensitive and high speed single photon detectors with a large cross-section and a low jitter. It also becomes apparent that the observed switching events are likely caused by constrictions (a local narrowing), which prevents devices to be biased at high enough currents, hence reaching high detection efficiencies. These issues could be resolved by optimization of the nanowire fabrication process. At present conditions, considering the photon flux on the detector area, we estimate the detection efficiency as $\sim 10^{-2}$%.

With the presently utilized deposition rate, the observed normal state resistivity in MgB$_2$ films increases sharply whereas the critical temperature drops down as the film becomes thinner than 5 nm. Nevertheless, it appears to be interesting to investigate if further reduction of the deposition rate would result in even thinner films (2–3 nm) with the desired degree of uniformity, low kinetic inductance, and high switch current density. Further steps shall also be taken towards increasing nanowire uniformity, i.e. avoiding constrictions and hence increasing the current density in the biased nanowire. We consider it important in order to reach a high quantum efficiency in SNSPDs made from MgB$_2$ films.

Acknowledgments

The authors would like to thank: Dr Olof Bäcke for conducting TEM analysis of MgB$_2$ samples; Professors X X Xi and K Chen for sharing their experience with the HPCVD process for MgB$_2$ films and enlightening discussions; and Usman Ul-Haq for the help with process development. MgB$_2$ film technology was developed under an ERC grant 308130-Teramix. This research was also supported by Swedish Research Council (Grant Nos. 2016-04189 and 2019-04345) and by Swedish National Space Agency (Grant No. 198/16). Devices were fabricated in the Nanofabrication Laboratory of Chalmers University of Technology.

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