Topical Review

High-\(T_c\) SQUID biomagnetometers

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Abstract

In this paper, we review the preparation technology, integration in measurement systems and tests of high-\(T_c\) superconducting quantum interference devices (SQUIDs) intended for biomagnetic applications. A focus is on developments specific to Forschungszentrum Jülich GmbH, Chalmers University of Technology, MedTech West, and the University of Gothenburg, while placing these results in the perspective of those achieved elsewhere. Sensor fabrication, including the deposition and structuring of epitaxial oxide heterostructures, materials for substrates, epitaxial bilayer buffers, bicrystal and step-edge Josephson junctions, and multilayer flux transformers are detailed. The properties of the epitaxial multilayer high-\(T_c\) direct current SQUID sensors, including their integration in measurement systems with special electronics and liquid nitrogen cryostats, are presented in the context of biomagnetic recording. Applications that include magnetic nanoparticle based molecular diagnostics, magnetocardiography, and magnetoencephalography are presented as showcases of high-\(T_c\) biomagnetic systems. We conclude by outlining future challenges.

Keywords: epitaxial heterostructures, high-\(T_c\) Josephson junctions, SQUIDs, biomagnetic measurement systems, magnetocardiography, magnetoencephalography, magnetic nanoparticles

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

1. Introduction

The field of biomagnetism encompasses the detection of extremely weak magnetic fields generated by biological systems, e.g., by ion currents in nerve cells and heart tissue or by magnetic nanoparticles (MNPs) of magnetite (Fe\(_3\)O\(_4\)) that occur naturally in living organisms due to bio mineralization or are used as a contrast agent for protein assays. Several applications of biomagnetism are relevant for non-invasive medical diagnostics, including the study of heart and brain function, peripheral nerve and spinal cord diagnostics, as well as the localization of cancerous tissue using processed MNPs as markers (see, for example, [1–3]).

Measurements of magnetic fields that are produced purely by living biological objects and originate from tiny ionic currents in the heart tissue or in neurons in the brain using magnetocardiography (MCG) and magnetoencephalography (MEG) provide valuable information about function, pathophysiology, and health conditions. Such measurements are, however, challenging. In order to have value in medical diagnostics and research, an MCG measurement should have
a signal-to-noise ratio (SNR) that is comparable with that of
electrocardiography of \(\sim 100\), which corresponds to a noise
equivalent magnetic induction (NEMI) in the case of MCG
that is lower than \(10 \, \text{fT Hz}^{-1/2}\) in the 1 Hz–1 kHz frequency
range. A similar sensitivity and frequency bandwidth are
required for sensors that are used in MEG to detect and
characterize deep sources of magnetic field in the human
brain. Sufficient sensitivity and frequency bandwidth for
biomagnetic measurements in the presence of relatively large
background magnetic fields of \(>100 \, \text{nT}\) can currently only be
achieved by using special sensors (‘biomagnetometers’) that
are based on superconducting quantum interference devices
(SQUIDs).

Another promising technique for neuromagnetic recording
utilizes atomic magnetometers (also referred to as optically-
pumped or spin exchange relaxation-free magnetometers). It has
been shown [4] that an optical-spectroscopy-based chip-
scale atomic magnetometer sensor with a magnetic sensitivity
of \(\sim 200 \, \text{fT Hz}^{-1/2}\) at 10 Hz is able to record spontaneous and
somatosensory evoked neuromagnetic fields. Johnson et al
used this technology to measure brain response from auditory
and median nerve stimulation [5]. Although atomic magnet-
ometers are promising because they do not require cooling,
a problem with the current technology is that the sensitivity
(\(\sim 15 \, \text{fT Hz}^{-1/2}\), dynamic range, and a too narrow frequency
bandwidth (\(\sim 10 \, \text{Hz}\)) of the sensors are not sufficient for pro-
cessing most MEG signals. Improvements have recently been
reported, where the authors claim that new atomic magnet-
ometers provide a NEMI of better than \(1 \, \text{fT Hz}^{-1/2}\) and do not
need to be shielded from the Earth’s magnetic field [6].
Theoretical studies [7, 8] have shown that atomic magnet-
ometers can provide a higher SNR and potentially replace
SQUIDs in MEG systems. A recent publication [9] also shows
a capability of neuromagnetic source localization using a single
atomic magnetometer (with a NEMI of \(15.9 \, \text{fT Hz}^{-1/2}\)) in the
2–80 Hz frequency range) placed sequentially at 13 locations.
However, an increase in the frequency range is an issue
because the sensitivity and frequency bandwidth of such a
sensor technology are mutually incompatible [10]. The issue of
cross-talk between sensors that are in close proximity to one
another will also be significant, due to either diffusion of the
gas from one laser readout beam to another or mutual induc-
tance between feedback coils surrounding the gas enclosures.
Furthermore, the utilization of atomic magnetometers for
neuromagnetic recording in less shielded environments is
limited by the dynamic range of the readout electronics. In
contrast, SQUIDs can be configured as gradiometric super-
conducting flux transformers that intrinsically remove environ-
mental noise during measurements by recording the gradient
of the magnetic flux. Moreover, flux counting readout elec-
tronics with a dynamic range of \(\sim 130 \, \text{dB}\) and a slew rate of
about \(5 \, \text{MHz}^{-1}\) can be used with low- and high-\(T_c\) direct
current (DC) SQUIDs [11].

Nearly all biomagnetometers in commercially available
MCG and MEG measurement systems rely on low-\(T_c\)
SQUIDs that operate at temperatures below 9 K and are
typically cooled with liquid helium. The use of liquid helium
as a coolant is inconvenient for medical personnel and results
in enormous operating costs, thereby preventing the wide-
spread distribution of such systems in ordinary hospitals. The
prospect of changing to liquid-nitrogen-cooled high-\(T_c\)
SQUIDs in future biomagnetometer systems would eliminate
the dependence on increasingly unreliable helium supplies
[12] and reduce risk for end users, while relaxing the demands
that are placed on them. It would also save a large fraction of
the operating costs. It is expected that the total cost (inves-
tment plus operating cost) of each high-\(T_c\) biomagnetometer
system would only be about half that of a corresponding
low-\(T_c\) system. When compared to a liquid helium system, the
liquid nitrogen cryostat design is far simpler, reducing the
overall system cost. The cryogen itself also has a much lower
evaporation rate and is plentiful, meaning that running costs
are far lower. As such, high-\(T_c\) SQUID technology may stim-
ulate the acceptance of biomagnetometer systems for more
widespread clinical use. High-\(T_c\) biomagnetometer systems
have the added advantage that cryostats with smaller separations between the superconducting sensor and the room
temperature environment of biologically active objects are
possible. The improved sensor-to-source proximity leads to
higher SNR and better representation and characterization of
the true source signal. For example, the dipole nature of
magnetic field sources in biological objects provides a boost in
signal power to high-\(T_c\) sensors that are within 1 mm of a
room temperature object. The SNR (e.g., in MEG recordings)
can then be high enough that a relatively large biomagnet-
ometer sensor NEMI of about \(50 \, \text{fT Hz}^{-1/2}\) at 77 K can be
acceptable [13–15]. However, state-of-the-art noise levels in
high-\(T_c\) technology (\(< 10 \, \text{fT Hz}^{-1/2}\) in the 1 Hz–1 kHz fre-
cquency range) are paramount for making biomagnetometer
systems that are not only cheaper and more accessible than
their predecessors, but also significantly better in terms of
providing relevant clinical diagnostic and monitoring infor-
mation and surgical guidance.

When compared to their low-\(T_c\) predecessors, high-\(T_c\)
biomagnetometers currently have limited clinical utility
because their technology is still too sophisticated and has yet
to reach the level of maturity that is needed for mass pro-
duction. The reasons for this situation are multifold. In con-
trast to low-\(T_c\) Nb technology, the most successful material
technology for high-\(T_c\) SQUIDs to date is \(\text{YBa}_2\text{Cu}_3\text{O}_7\)–\(x\)
(YBCO), a ceramic oxide superconductor that has intrinsic
complications, including a much shorter coherence length,
d-wave symmetry of the superconducting order parameter,
and relatively weak pinning forces at liquid nitrogen tem-
perature. All of these issues demand very different technolo-
gies for the preparation of films, Josephson junctions (JJs),
and thin film multilayer heterostructures, when compared to
low-\(T_c\) technology. High-\(T_c\) YBCO SQUIDs must be made
from high quality thin film epitaxial heterostructures in an
oxygen atmosphere at a relatively high substrate temperature
of \(< 800 \, ^\circ \text{C}\). Currently, the best high-\(T_c\) sensor technology
consists of DC SQUIDs that are coupled to thin film super-
conducting multilayer flux transformers. Their fabrication is
so challenging that only a handful of dedicated research
groups have managed to produce high-\(T_c\) SQUIDs that have
white noise levels of below \(10 \, \text{fT Hz}^{-1/2}\) (at 77 K) at

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frequencies of above a few Hz [16–20]. The main challenge is to avoid excessive 1/f low frequency noise in the high-$T_c$ superconducting thin film multilayer flux transformer, which can originate from the presence of parasitic grain boundaries (GBs) in the top epitaxial YBCO film, especially near the edges of the underlying layers, and/or due to degraded superconducting parameters in the bottom YBCO film.

In this paper, we provide an overview of the sensor technology, measurement system construction, and examples of applications of currently available high-$T_c$ biomagnetometers. We outline the deposition and structuring of the epitaxial oxide heterostructures, materials for substrates, epitaxial bilayer buffers and multilayer flux transformers, and step-edge and bicrystal JJs. We also present the layout and properties of epitaxial multilayer high-$T_c$ DC SQUID sensors, as well as their integration in measurement systems with special electronics and liquid nitrogen cryostats for different applications. Future challenges in the field are then addressed.

2. Experimental technique

In this section, the technology and related properties of high-$T_c$ DC SQUID biomagnetometer sensors, their integration in high-$T_c$ biomagnetic measurement systems with liquid nitrogen cryostats and SQUID readout electronics are described. High-$T_c$ DC SQUID sensors were separately produced at Forschungszentrum Jülich GmbH (Jülich) and at Chalmers University of Technology (Chalmers) using different technologies. The Jülich technology includes high-oxygen-pressure sputtering [21] for the deposition of YBCO films, step-edge JJs [22] for high-$T_c$ DC SQUIDs and superconducting multilayer thin-film flux transformers with the input coil inductively coupled to the SQUID loop in a flip-chip configuration. At Chalmers, laser ablation for the deposition of YBCO films, bicrystal JJs [23] for DC SQUIDs and direct-coupled high-$T_c$ DC SQUIDs (with the pickup loop made on the same substrate as the SQUID and galvanically connected to the SQUID loop) were used. Jülich used modulation DC SQUID electronics and cryostats, which were developed at the Kotel’nikov Institute of Radio Engineering and Electronics RAS and produced at Cryoton Co. Ltd [24], while Chalmers used direct amplification electronics from Magnicon and cryostats from ILK Dresden [25].

2.1. High-$T_c$ DC SQUID biomagnetometer sensors

Bicrystal JJs are one of the earliest technological developments in high-$T_c$ DC SQUID fabrication [23]. The process is advantageous because it is simple and reproducible, although it suffers from a lack of readily-available high quality bicrystal substrates, as well as limited design freedom imposed by the fact that the placement of JJs is constrained to the bicrystal boundary.

The Chalmers bicrystal high-$T_c$ DC SQUID biomagnetometer sensors are made on 1 mm thick bicrystal SrTiO$_3$ (STO) substrates with a 24° misorientation angle. An ultrahigh-vacuum pulsed laser deposition system (DCA instruments) is used to deposit 200–300 nm thick YBCO films on the bicrystal substrates. A seed layer of between 20 and 50 nm of CeO$_2$ may be included for improving the epitaxy of the YBCO. During YBCO deposition, the oxygen pressure is maintained at 0.5–1.0 mbar. The sample is annealed over ~2 h at a temperature of 500°C–600°C, in order to ensure proper oxygen doping. An ion beam etching (IBE) and standard photolithography are used to pattern single layer bicrystal high-$T_c$ DC SQUID biomagnetometers. Further details about film deposition, design, and fabrication can be found elsewhere [26, 27].

Each of the Jülich step-edge high-$T_c$ DC SQUID biomagnetometer sensors was made on two separate 1 mm thick single crystal (001) oriented MgO substrates: one 10 mm × 10 mm substrate for two SQUIDs and one 30 mm diameter wafer for a superconducting flux transformer. The low cost and good availability of single crystal MgO substrates, a thermal expansion coefficient of $\sim 13.4 \times 10^{-6}$ K$^{-1}$ that matches well with YBCO, a low dielectric constant ($\varepsilon \approx 10$) and low microwave losses ($\delta \approx 10^{-5}$) are all decisive advantages of using MgO as a substrate material for the preparation of heterostructures with YBCO. The lattice mismatch between YBCO and MgO is ~9% for a cube-on-cube mutual orientation. Although this mismatch results in difficulties for the cube-on-cube epitaxial growth of YBCO films on MgO, it facilitates growth on other crystallographic planes and grahopheptaxial growth. We take advantage of the fact that an YBCO film grows c-axis-oriented on 45° tilted MgO substrates [28], as well as on the surfaces of 45° tilted step edges on (001) MgO substrates [20]. At interfaces/step-edges between these crystallographic orientations on the substrate, c-axis-oriented layers of YBCO consistently form two symmetrical GBs with an exact 45° [100]-tilted misorientation angle between them. Surprisingly, this situation can be achieved despite slight deviations in the slope angle of the step edges from an exact 45° value [20]. High quality and relatively smooth (droplet-free) YBCO films are deposited by high oxygen pressure sputtering in pure oxygen (99.999%) at a pressure of ~3 mbar [21].

An IBE and the negative photoresist nLOF2020 were used for the preparation of 45° steps on (001) MgO substrates (see the scanning electron microscopy (SEM) images in [29]). A second ion beam milling step was used to clean the surface [20, 30], in order to remove residual resputtered material, including undesirable ‘fences’ that appear when resputtered material gathers at the top corners of the steps during the first ion beam milling procedure. A much longer ion beam milling step was additionally used to texture the surface of the MgO substrate in the form of linear trenches along the [100] and [010] directions [20, 22, 29]. The bottom corner of the step was smoothed due to the simultaneous effect of the angular dependence of ion beam milling and the partial redeposition of etched material.

The 45° slope of the step edges on MgO is gentle enough for the homogeneous deposition of thin film buffer layers by high oxygen pressure sputtering. The homogeneity and quality of the superconducting YBCO films was improved significantly by the introduction of a double buffer...
layer: a bottom 10 nm thick non-superconducting YBCO film as a ‘seed’ layer that was covered by a 30 nm thick SrTiO$_3$ (STO) film as a ‘blocking’ layer [29]. Figure 1 shows an SEM image of a YBCO film deposited on such a double-layer-buffered 45° step on an MgO substrate, as well as a high-resolution transmission electron microscopy (HRTEM) image of the 45° [100]-tilted GB in the top (superconducting) YBCO layer at the lower corner of the step. The top YBCO layer had a thickness of ~200 nm, a $T_c$ of ~93 K and a critical current density of ~5 MA cm$^{-2}$ at 77 K. We protected the superconducting film with a 5 nm thick Pt film that passivated the junctions and provided a better electrical contact for bonding.

The surface texturing changes the in-plane orientation of the YBCO film on the MgO substrate and step edges. SEM images of the surface of the YBCO film in the step-edge area show rectangular growth spirals with 1 nm high steps between flat areas [31]. The orientations of such spirals indicate that all of the grains in the YBCO film are aligned in-plane, with their $a$- or $b$-axes normal to the corners of the step. In this case, the superconducting critical current density is maximized and constant along both GBs. Such texturing of the MgO surface also helps to increase the relevant JJ parameters, reduce the meandering of GBs, and improve the overall characteristics of the SQUIDs.

High-$T_c$ JJs are made by patterning bridges of width $w$ ~ 2 $\mu$m in the YBCO-STO-YBCO heterostructure across the substrate step. Both the upper and the lower GBs in each step-edge junction have similar cross-sections and misorientation angles and, consequently, similar critical currents and normal state resistances $R_n$. They are connected serially and act together synchronously as a single JJ with twice the normal state resistance and half of the capacitance of a single GB. For such an effective JJ, a total capacitance $C$ of ~10 fF, a normal state resistance $R_n$ of ~20 $\Omega$, and a characteristic voltage $V_c = I_c R_n$ of ~800 $\mu$V at 77 K were obtained from the analysis of the $I(V)$ characteristics of the junction. A capacitance $C$ of ~10 fF of step edge JJs on an MgO substrate was reported previously [32]. Such a low capacitance of a high-$T_c$ JJ is advantageous for lowering the intrinsic flux noise $S_\Phi$ of high-$T_c$ DC SQUIDs made with such JJs [33, 34]:

$$S_\Phi \approx 32 \kappa_b T S (L_s C / \beta c)^{1/2},$$

where $L_s$ is the SQUID loop inductance, $\beta c = 2 \pi I_c R_s^2 C / \Phi_0 \approx 0.4$ is the Stewart–McCumber parameter of the JJ and the magnetic flux quantum $\Phi_0 = h/2e = 2.068 \times 10^{-15}$ T m$^2$, where $h$ is the Planck constant and $e$ is the electron charge. The Stewart–McCumber parameter remains almost constant for high-$T_c$ JJs of different width and/or cross-sectional area, whereas in the case of low-$T_c$ JJs the normal state resistance $R_n$ is determined by the shunt resistor and is independent of the JJ area.

Comparing the intrinsic flux noise of low-$T_c$ and high-$T_c$ SQUIDs is illustrative of other benefits that are inherent to high-$T_c$ technology. For example, the far lower operating temperatures of low-$T_c$ SQUIDs should reduce their intrinsic noise levels $S_\Phi$ when compared to high-$T_c$ technology by a factor of $77/4.2 \approx 20$. This factor is, however, almost compensated by the ~100 × higher capacitance of Nb-based planar JJs (~1 pF), resulting in comparable NEMI values of ~5 $\mu$T Hz$^{-1/2}$ for high-$T_c$ and low-$T_c$ SQUIDs, in spite of the much higher operating temperature of high-$T_c$ SQUIDs.

For applications of DC SQUIDs, the spread in JJ critical current should be as low as possible, and preferably below 10%. This spread depends on the morphologies of the GBs and on the precision of the patterning of the YBCO films. 45° [100]-tilted GBs on step-edge JJs on MgO are very straight: their deviation is <~100 nm, which is significantly smaller than the ~200 nm deviation of the 24° [001]-tilted GBs in bicrystal-substrate-based JJs [19]. More than 100 step-edge JJs with 45° step edges on MgO substrates were prepared during optimization of the technology. The observed spreads in critical current $\delta I_c / I_c$ and $I_c R_n$ product depend on the microstructural quality of the JJs. Without using an STO blocking layer, typical values for $\delta I_c / I_c$ of ~20% and for the $I_c R_n$ product of ~300 $\mu$V were measured at 77 K [20]. Meandering of the GBs and mutual misorientation of grains in the YBCO film at the step edge area were reduced by properly texturing the MgO substrates after step preparation. The STO
blocking layer has a reduced density of holes and precipitates in the YBCO film. As a result, the spread in critical current \( I_{c}/I_{0} \) was reduced to \( \sim 10\% \) [29] and values of the \( I_{c}R_{m} \) product of up to \( \sim 800 \, \mu\text{V} \) were observed at 77 K. As a comparison, for bicrystal-substrate-based JJs values of the \( I_{c}R_{m} \) product of up to \( \sim 400 \, \mu\text{V} \) at 77 K and \( \sim 1 \, \text{mV} \) at 63 K were achieved [19]. At present, the spread in critical current \( I_{c}/I_{0} \) in 2 \( \mu \text{m} \) wide JJs is determined primarily by the precision of the patterning of the YBCO films of \( \sim 0.2 \, \mu\text{m} \) using deep-UV photolithography. This value can be improved in the future by using electron beam lithography. Indeed, the most reproducible patterning of YBCO films with variations in width \( \delta w \) of \( <0.2 \, \mu\text{m} \) is achieved using electron beam lithography, a hard carbon film mask, and IBE (see for example [35] and references therein).

The operational principle of a SQUID is based on its sensitivity to changes in magnetic flux passing through its loop. The larger the SQUID loop, the more flux penetrates it for a fixed magnetic field magnitude, and thus the higher its sensitivity. However, this relationship is limited by equation (1); a larger SQUID loop leads to higher inductance and therefore lower sensitivity. Furthermore, for design reasons, a screening parameter \( \beta_{L} = 2L_{S}/\Phi_{0} \sim 1 \) should be chosen, which is a compromise between a sufficient modulation depth and a large enough value of \( L_{S} \) to couple flux. For a high-\( T_{c} \) SQUID operating at 77 K, the critical current (i.e., the Josephson coupling energy) needs to be higher than 16 \( \mu\text{A} \) to avoid thermal smearing and this puts a limit on the SQUID inductance \( L_{S} \sim 60–100\,\text{pH} \). A tradeoff is therefore made, whereby the SQUID loop is kept small \( (\sim 10^{-5} \, \text{cm}^2) \) and a superconducting flux transformer is used to collect magnetic flux from a relatively large area pickup loop (e.g., about 3 \( \text{cm}^2 \)) and to transfer it inductively to the SQUID loop via a multiturn input coil or by direct injection of the induced current into the SQUID loop using a galvanic connection between the pickup loop and the SQUID loop.

Two types of flux transformer are referred to as ‘direct-coupled’ and ‘flip-chip’. Direct-coupled sensors are simple to fabricate because the flux transformer can be defined together with the SQUID in a single lithography step, as is the case for single-layer sensors. Fabrication challenges associated with multilayer technology in YBCO mean that the flux transformer is typically fabricated on a second substrate that is glued to the SQUID substrate in a ‘flip-chip’ configuration. Despite this complication, flip-chip magnetometers are advantageous because they have much better sensitivity when compared to direct-coupled magnetometers. The effective area \( A_{\text{eff}} \) is \( \sim 10 \times \) larger for flip-chip designs, when compared to equivalent direct-coupled designs in the case of 20 \( \mu\text{m} \) pickup coils. This gain increases as a function of the size of the pickup coil because direct-coupled sensors suffer from an increasing inductance mismatch between the large pickup coil and the SQUID. This mismatch can be mitigated in the more flexible flip-chip design by careful selection of flux transformers with inductances of the pickup coil and input coil that are approximately equal to each other. The effective area of such a flip-chip magnetometer \( A_{\text{eff}} \sim A_{\text{pu}} \sqrt{L_{\text{pu}}} \sim a^{3/2} \), where \( A_{\text{pu}} \) is the area, \( L_{\text{pu}} \) is the inductance, and ‘\( a \)’ is an approximate measure of the size (i.e., the diameter of a circular loop or the side dimension of a square loop) of the pick-up loop of the flux transformer. The flip-chip approach is currently the only method that has achieved the required values of NEMI of below 10 \( \text{fT Hz}^{-1/2} \) at 10 Hz and 77 K. Our high performance high-\( T_{c} \) superconducting multi-layer flux transformer is made from epitaxial thin-film heterostructures that contain two high-\( T_{c} \) superconducting layers of YBCO separated by an insulating layer, which consists of \( \text{PrBa}_{2}\text{CuO}_{x} \) (PBCO) and STO thin films [36]. A critical process that is needed for the preparation of high-\( T_{c} \) superconducting multilayer thin film flux transformers is based on chemical wet etching in Br-ethanol solution [37]. This step is used to pattern the bottom YBCO film, while preventing issues that are critical for multilayer epitaxy and unavoidable when IBE is used; the creation of ‘fences’ at the edges of the photoresist mask and ion-beam-induced damage to the substrate surface. Appropriate superconducting parameters of all of the superconducting films are a prerequisite for the preparation of low-noise biomagnetometers, especially in the low frequency range. GBs should be avoided in the top YBCO layer because they cause increased \( 1/f \) noise of the sensor [38]. Chemical etching results in gently sloping edges of the patterned structures, which are advantageous for avoiding GBs in the top YBCO layer [37]. The input coil includes a \( \sim 200 \, \text{nm} \) bottom YBCO film that is covered by a \( \sim 400 \, \text{nm} \) thick layer of PBCO-STO insulator and a 600–1000 nm top YBCO film. The dynamic range of the flux transformer is limited by the critical current of the input coil, which is proportional to the thickness of the top YBCO film. It should be noted that oxygenation of the bottom YBCO layer should be performed despite the presence of the STO insulator layer deposited above it. Fortunately, STO enables sufficient diffusion of oxygen ions for oxygenation of the bottom YBCO film at a substrate temperature of \( \sim 500 \, ^{\circ}\text{C} \). The empirically-obtained time required for the full oxygenation of a bottom YBCO film covered by a 0.5 \( \mu\text{m} \) thick STO film is \( \sim 2 \, \text{h} \) at 500 \( ^{\circ}\text{C} \). The gently sloping edges of patterned structures and full oxygenation of the YBCO films have helped to avoid increased \( 1/f \) noise of the superconducting multilayer flux transformers.

A schematic diagram of the SQUID flip-chip magnetometer with coupling to control electronics and a schematic 3D layout demonstrating the flip-chip arrangement of the SQUID, flux transformer and modulation coil are shown in figures 2(a) and (b), respectively. The SQUID has a 1 \( \text{mm} \) washer and a 3 \( \text{mm} \) direct-coupled pick-up loop. The 14-turn input coil of the flux transformer is inductively coupled to the SQUID washer, while the direct-coupled loop is inductively coupled to the modulation/feedback coil of the SQUID readout electronics and also serves as one of the superconducting electrodes for current biasing of the SQUID.

High-\( T_{c} \) superconducting flux transformers with multi-turn input coils can be made on STO, \( \text{LaAlO}_{3} \) (LAO), or MgO substrates. As for SQUID fabrication, MgO wafers have the advantage of a well-matched thermal expansion coefficient to YBCO. Sufficiently thick YBCO films, such as those typically used for the second superconducting layers of flux
transformers, develop cracks when they are deposited on STO and LAO substrates, whereas no cracks are observed for YBCO films that are as thick as 5 \( \mu \)m and are grown on MgO substrates [40]. A noteworthy limitation for the use of MgO substrates is that they require epitaxial double-layer buffering.

The preferred seed layer for the preparation of flux transformers is BaZrO\(_3\) (BZO) because its lattice constant is similar to that of MgO, its crystal structure is similar to that of the second (STO) buffer layer that is destined to grow on top of it, and it is insensitive to the Br-ethanol solution that is used for patterning the bottom YBCO layer. This epitaxial BZO–STO buffer layer also protects the hygroscopic surface of the MgO substrate against degradation in air and during lithographic procedures. Whereas antiphase boundaries at the BZO–MgO interface have been observed to spread through the STO layer, they usually disappear at the STO–BZO interface [41]. The STO layer initially grows with an expanded lattice constant to match that of the underlying BZO. However, within a few unit cells of the STO–BZO interface the STO lattice constant relaxes to its bulk value. The surface of the STO layer on which the top YBCO layer grows therefore has a lattice constant and a microstructural quality that are similar to those of a single-crystal STO substrate. However, the overall thermal expansion coefficient is still determined by the 1 mm thick MgO wafer.

Figure 3 shows an example of the noise spectral density of a 16 mm high-\( T_c \) DC SQUID flip-chip magnetometer with step-edge junctions measured inside shields consisting of three layers of \( \mu \)-metal and one of a high-\( T_c \) superconductor. The effective area \( A_{\text{eff}} \) and the flux-to-field transformation coefficient of this magnetometer are approximately 4.6 mm\(^2\) and 0.45 nT/\( \Phi_0 \), respectively. In the white noise region above \( \sim 10 \) Hz, the magnetic flux noise \( \sqrt{S_\Phi} \) and NEMI are approximately 10 nT/\( \Phi_0 \) Hz\(^{-1/2}\) and 5 fT Hz\(^{-1/2}\), respectively, at 77 K. The NEMI \( B_N \) can be estimated based on the properties of the SQUID, the flux transformer, and the coupling between the two, according to the equation

\[
B_N = \frac{L_{\text{pu}} + L_1}{kA_{\text{pu}} \sqrt{L_1L_S}} S_\Phi^{1/2},
\]

where \( L_S \approx 100 \) pH and \( L_1 \approx L_{\text{pu}} \approx 40 \) nH are the inductances of the SQUID loop and the input coil, respectively, while \( k \approx 0.7 \) is the coupling coefficient and \( A_{\text{pu}} \approx 256 \) mm\(^2\) is the area of the pickup loop. Below \( \sim 10 \) Hz, additional noise in experimental recordings may be generated by the biomagnetometer itself and external sources, e.g. Abrikosov vortex creep in the superconducting shield or residual laboratory environmental noise coupling via electrical connections. Increased low frequency noise of the SQUID can also be a first indication of ageing in one of the superconducting films of the SQUID and/or superconducting multilayer flux transformer.

Ageing is a significant problem for YBCO films, as their performance is highly dependent upon their oxygen doping and chemistry, both of which are extremely sensitive to the environment. The degradation of the superconducting parameters of YBCO films can be prevented by ensuring vacuum-tight encapsulation of the magnetometers. An example of such a capsule is shown in figure 4. Encapsulated sensors have lower intrinsic noise and longer-term stability because the vacuum-tight capsule dampens external thermal fluctuations, isolates them from ambient air and humidity, and provides mechanical protection of the sensors, thereby ensuring a long and reliable service life. This is especially important for delicate flip-chip sensors with multilayer flux transformers. Encapsulation also significantly simplifies their
2.2. Cooling of high-$T_c$ DC SQUID biomagnetometers

High-$T_c$ biomagnetometers operate at $T \sim 77$ K, which corresponds to an optimal value of the Stewart–McCumber parameter $\beta_c \sim 0.4$ of the bicrystal and step-edge JJs. The cheapest and most convenient cooling method is to simply immerse the sensors in liquid nitrogen at normal atmospheric pressure. The price of liquid nitrogen is presently $\sim 300$ times lower than that of liquid helium ($\sim 15$ €/l for liquid helium and $\sim 0.06$ €/l for liquid nitrogen). This difference in price is expected to grow, as helium is a finite resource, whereas nitrogen comprises $\sim 80\%$ of the atmosphere. Liquid nitrogen is also $\sim 60$ times more efficient in terms of cooling power. The heats of vaporization of helium and nitrogen are $2.68\,\text{J/l}$ and $161\,\text{J/l}$, respectively. High-$T_c$ biomagnetometers can therefore be cooled in cryostats that have fewer radiation shields and can be placed closer to the outer wall/room temperature environment than low-$T_c$ biomagnetometers. In a standard low-$T_c$ MEG system, the spacing between the pickup loop of the sensors and the room temperature cryostat is typically $\sim 20$ mm. For a high-$T_c$ system with a similar cryogen consumption (boil-off rate), this spacing can be below $\sim 5$ mm. From a practical perspective, the hold time of a liquid nitrogen cryostat is on average about five times longer than that of a liquid helium cryostat. This difference promises to increase the availability of high-$T_c$ biomagnetometer systems when compared to low-$T_c$ technology, while reducing the person-hours needed to maintain them. For example, a 50 l liquid-nitrogen-based full-head MEG system can run continuously for more than a month without being refilled. In contrast, the time period is $\sim 1$ week for a 100 l liquid-helium-based MEG system that is presently state-of-the-art.

The relaxed cooling requirements of high-$T_c$ biomagnetometer technology result in an additional benefit of a reduced spacing between the sensor and the object, i.e., the biomagnetic source. A smaller distance between the biomagnetometer and the object leads to improved signal levels arising from dipolar-like sources that generate the biomagnetic fields. The magnitude of the magnetic field generated by current dipoles, such as those associated with neuromagnetic activity recorded in MEG, decays as $\sim 1/r^2$ with distance $r$ between the biomagnetometer and the object. For magnetic dipoles, such as those associated with MNPs, the equivalent dependence is $\sim 1/r^3$. As such, a reduction in spacing, $r$, by a factor of 3 (which is reasonable when comparing cryostats for high-$T_c$ and low-$T_c$ SQUIDs) leads to an order of magnitude gain in signal magnitude. For similar NEMI values and sensor sizes, the SNR obtained during the detection of superficial sources of biomagnetic signals by a high-$T_c$ biomagnetometer system can therefore be significantly higher than that for a low-$T_c$ biomagnetometer system. The spatial resolution can also be enhanced because biomagnetic fields can have higher spatial frequencies at an object surface. For example, in the case of MEG, 40% more information can be obtained from the human brain by using a high-$T_c$ MEG system when compared to the state-of-the-art in a low-$T_c$ MEG system [14]. If enhanced signal amplitude of the high-$T_c$ biomagnetometer system does not benefit a particular

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Figure 4. Photograph of an encapsulated high-$T_c$ biomagnetometer that includes a superconducting flux transformer with a round pick-up loop, whose outer diameter is 20 mm. The button-shaped fiberglass capsule has an outer diameter of $\sim 24$ mm and a total thickness of $\sim 10$ mm.

Chalmers has produced ‘directly-coupled’ high-$T_c$ biomagnetometers on 10 mm bicrystal STO substrates that have an in-plane misorientation angle of 24° [13]. The preparation of such sensors includes only a single layer of YBCO and is much simpler when compared to the multilayer technology described above. However, the typical NEMI (of about $50\,\text{fT}\,\text{Hz}^{-1/2}$ at 10 Hz and 77 K) is much higher than the NEMI of flip-chip magnetometers. The higher NEMI values impose restrictions on the construction and possible applications of biomagnetic measurement systems. In order to achieve comparable SNR values for the same measurement time, the sensor should be placed much closer to the object in the case of MEG measurements. However, for the recording of MCG and magnetic signals originating from MNPs, a NEMI value of $50\,\text{fT}\,\text{Hz}^{-1/2}$ at 10 Hz and 77 K appears to be sufficient.
behind the thin room temperature sapphire window of the cryostat. A sapphire rod that is thermally anchored to the liquid nitrogen bath is the magnetometer used for MEG recordings, as it sits on top of the Institute biomagnetometer in the ILK Dresden cryostat at the Karolinska.

Figure 5. The cryostat was developed and produced by ILK Dresden (© 1969 IEEE. Adapted, with permission, from [15]). (a) Cutaway schematic diagram of the cryostat. (c) Photograph of the ‘directly-coupled’ high-$T_c$ DC SQUID magnetometer used for MEG recordings, as it sits on top of the sapphire rod that is thermally anchored to the liquid nitrogen bath behind the thin room temperature sapphire window of the cryostat.

application, then it can be balanced by a reduction in the total system cost by using less sensitive SQUIDs and/or reducing the magnetic shielding.

The first tests of the Chalmers ‘directly-coupled’ high-$T_c$ biomagnetometers for magnetic immunoassays (MIAs) and MEG were performed using a fiberglass-reinforced epoxy cryostat with a 1 l liquid nitrogen tank and a hold time at the base temperature $T \sim 73$ K of approximately 8 h, as shown in figure 5. The cryostat was developed and produced by ILK Dresden [25]. We pump on the liquid nitrogen bath to reduce its boiling temperature and to achieve an operating temperature below 77 K. A thermal link connects the liquid nitrogen bath to a sapphire rod, on which the SQUID chip is mounted, such that the cryostat can be operated both in the standard ‘sensor down’ and at arbitrary, including ‘sensor up’, tilt angles. Furthermore, the sensor is housed in vacuum behind a thin (200 μm) sapphire window at room temperature. In order to reduce the thermal load of ambient radiation, the window can have a thin layer of reflective Au deposited on its backside. However, the Au should be patterned in the form of a 5 mm grid to limit the conducting paths of eddy currents that could otherwise contribute to increased values of NEMI. Micrometer screws support the inner liquid nitrogen + sensor structure so that it can be raised and lowered with respect to the room temperature window. Careful adjustment of the screws while the cryostat is cold allows the high-$T_c$ biomagnetometer to be placed within less than 1 mm of the room temperature environment, while maintaining its operating temperature.

First tests of Jülich ‘flip-chip’ high-$T_c$ biomagnetometers for MCG and MEG were performed using a fiberglass cryostat with a 1.5 l liquid nitrogen tank and a hold time of 10 d. The cryostat was developed and produced by Cryoton Co. Ltd [24]. The cold–warm distance at the cryostat bottom is about 10 mm. Two encapsulated 16 mm high-$T_c$ biomagnetometers with NEMI values of $\sim 5 \text{ fT Hz}^{-1/2}$ at 77 K were fixed ‘on-axis’ parallel to each other on a fiberglass holder and immersed in liquid nitrogen, as shown in figure 6. The electronic gradiometer baseline of 10 cm is determined by the distance between the flux transformers of the magnetometers. The bottom sensor is intended for detection of the signal from the object, while the top magnetometer serves as a reference, e.g., for reducing environmental noise. The distance between the pickup loop of the bottom magnetometer and the outer surface of the cryostat is about 15 mm. Both the ILK Dresden and the Cryoton single (signal) channel cryostats are convenient for sensor development proof-of-principle tests for different applications. However, they should be replaced by a multi-channel system, e.g., for practical use in MCG or MEG.

The use of a multichannel biomagnetometer system for MCG and MEG saves measurement time and also allows better localization and characterization of sources of magnetic field in the heart and brain, respectively. The sensors should be placed in a dense array and located as close as possible to the chest/scalp and to neighboring sensors. Cross-talk measurements [39] have confirmed the possibility of building such closely-packed arrays of high-$T_c$ biomagnetometers. Encapsulated high-$T_c$ biomagnetometers are sufficiently small that more than 100 sensors can be placed above and around an adult human head in the form of a single helmet-shaped cryostat. One limitation to such an approach is that the variety of individuals’ head sizes and shapes requires varying the spacing between the sensors and the scalp and thus reduced proximity to some brain regions. This problem is inherent to state-of-the-art MEG systems, but can be overcome by using high-$T_c$ biomagnetometer systems that are designed with a mechanically flexible set of cooling systems that can be adjusted to maintain close proximity between each sensor and the scalp.

In principle, each channel can be placed in an individual cryostat that has a small area at its lower end and is adjusted individually to the scalp [42]. The advantage of such a
segmented helmet construction is the possibility to realize a SQUID-to-scalp separation down to <3 mm by using a very thin wall at the bottom end of a small cryostat. The main disadvantage is the increased tangential separation between the channels due to the sidewalls of the cryostat, which can lead to attenuation of high spatial frequencies in the MEG signals. However, cryogen transfer and cryostat costs are less critical for high-$T_c$ systems when compared to low-$T_c$ systems, as suggested in [42]. Cryogen-free cooling systems (see, e.g., [43]) can also be used, provided that their magnetic noise is sufficiently low.

High-$T_c$ biomagnetometers may be placed very close (at a distance of <3 mm) to the scalp by fixing them in the vacuum space of a cryostat inside thermal conducting sockets and cooling them sideways, for example with the help of vacuum grease. Alternatively, the sensors can be placed in liquid nitrogen inside individual dimples on the inner side of the cold wall.

2.3. Readout electronics for high-$T_c$ DC SQUID biomagnetometers

At an optimal bias current, the SQUID voltage $V$ is periodic as a function of magnetic flux $\Phi$ through the loop of a DC SQUID, with a period given by the flux quantum $\Phi_0$. Linearization of the $V(\Phi)$ transfer function is provided by SQUID readout electronics operating in a flux-locked loop mode using a modulation scheme [44] or direct amplification [45]. The primary advantage of the modulation scheme is an increase in the readout voltage signal that is available to the field effect transistor in the first amplification stage of the SQUID readout electronics. This is caused by a boost in effective SQUID output impedance by the application of a matching transformer that serves as a low-noise preamplifier. The primary disadvantage of the modulation scheme is that the bandwidth, which is typically $\sim 20$ kHz, is limited by the lock-in demodulation circuit. However, because most relevant biomagnetic signals do not contain useful information above a few kHz, a 20 kHz sensor bandwidth is sufficient for most biomagnetic applications.

The critical currents and normal resistances of JJs constituting DC SQUIDs have low frequency fluctuations with a spectral density $S \sim 1/f$. In the case of low-$T_c$ DC SQUIDs with Nb–Al$_2$O$_3$–Nb tunnel JJs, these fluctuations are weak and barely affect the SQUID low frequency noise [46]. On the other hand, these fluctuations are the dominant source of $1/f$ noise in high-$T_c$ DC SQUIDs [47]. In-phase fluctuations of the critical current are eliminated by using the ‘standard’ flux modulation scheme with negative feedback. Readout electronics with an additional bias-reversal modulation scheme intended for the reduction of $1/f$ noise originating from out-of-phase critical-current fluctuations were proposed a long time ago (see [47–49]) and have been included in commercial SQUID electronics units (from, e.g., Tristan Technologies, Magnicon, EZ SQUID, Supracon, STAR Cryoelectronics and Cryoton). The bias reversal circuit simultaneously switches the polarities of the bias current and the magnetic flux, thereby effectively canceling the out-of-phase critical-current fluctuations, while the system remains sensitive to changes in the $V(\Phi)$ curve due to the external magnetic flux. Bias reversal effectively reduces the ‘knee frequency’ (at which the $1/f$ and white noise contributions to the power spectral density are statistically equal) from a few kHz to a few Hz.

A special high-$T_c$ bias reversal DC SQUID readout electronics system that is used in the ‘flip-chip’ biomagnetometer experiments described below was developed at the Kotel’nikov Institute of Radio Engineering and Electronics RAS and produced by Cryoton Co. Ltd [24, 50]. The modulation and bias-reversal signals are formed in each channel by a programmable logic device. Furthermore, the implementation of correct compensating signals and the removal of transient voltage peaks generated by parasitic inductions during switching of bias current are allowed. In this high-$T_c$ SQUID readout electronics system, the current-bias-reversal and modulation signals are rectangular, with a frequency of the modulation signal that is twice that of the current-bias-reversal. Two special compensation signals help to remove the residuals of the current-bias-reversal in the primary winding of the matching transformer and to compensate, e.g.,

![Figure 6](image)
the slight asymmetry in the critical currents of the JJs in a given SQUID. In order to reduce crosstalk between the SQUID biomagnetometers, both the main and the bias-reversal modulations and feedback should be applied locally to the SQUID by a coil that has a much smaller outer size when compared to the size of the pickup loop [39]. Direct injection of current into the SQUID loop [51, 52] is another possibility for generating a locally applied modulation-feedback signal, which is convenient for single-chip integrated DC SQUID magnetometers.

3. Applicability test for high-$T_c$ DC SQUID biomagnetometers

In this section, a primary focus is placed on experiments that demonstrate the applicability of high-$T_c$ DC SQUID biomagnetometers for MCG, MEG, and MIA measurements performed in [13, 15, 20, 53–55].

For ‘direct-coupled’ biomagnetometers, all recordings were carried out using ILK Dresden cryostats and Magnicon DC SQUID electronics SEL-1 with AC bias reversal. The magnetometers that were used for MEG recordings had 8 mm pickup loops. Recordings were performed in 2-layer magnetically shielded rooms (MSRs), one of which is located at Imego AB (now part of RISE Acreo) in Gothenburg and the other at the Karolinska Institute’s NatMEG lab in Stockholm. In the case of MIA, a planar gradiometer was used with a figure-of-eight design and a 3 mm baseline. MIA recordings were performed in a simpler electrically-shielded room. This work was performed using either single- or independent two-channel systems.

MCG and MEG measurements with ‘flip chip’ biomagnetometers were performed using a two-channel system with a signal and a reference channel, each of which were high-$T_c$ DC SQUID biomagnetometers with 16 mm pickup loops. A two-channel bias-reversal DC SQUID electronics (Cryton) unit enabled simultaneous operation of the magnetometers. The experiments were performed in standard MSRs in the Franz-Volhard Hospital (MCG measurements [53]) and the Institute of Neurosciences and Medicine, Forschungszentrum Jülich (MEG measurements [20, 54]). The shielding factor of the MSRs was about $10^4$ at 50 Hz and about 10 at 0.1 Hz. The residual magnetic flux density in the middle of the MSRs was $\sim 100$ nT. A slow variation of the magnetic field in the middle of the MSRs was observed during daytime. The typical time constant of this drift was $\sim 10$ s with an amplitude of up to 5 nT. No active compensation of this low frequency environmental field noise was employed during the measurements. The feedback of the electronics was adjusted to follow the maximal changes of the magnetic field.

3.1. Application of high-$T_c$ DC SQUID biomagnetometers for MCG

Cardiovascular diseases are the most common cause of early disability and death globally, contributing to about 17.3 million deaths/year (30% of all deaths in the year 2008), with a prognosis that this number will grow to more than 23.6 million deaths/year by 2030 [56]. The early detection of heart diseases has enormous potential benefits across the world because it opens up important possibilities to prevent or even reverse disease progression and to reduce its devastating impact on later life. The most common non-invasive test that is used to diagnose cardiovascular diseases is the electrocardiogram (ECG), which is $\sim 50\%$ accurate in the diagnosis of the leading contributor to cardiovascular disease: myocardial ischemia. Although MCG is rarely used, it is a non-invasive and contact-free technique that has impressive sensitivity ($\sim 91\%$) and specificity ($\sim 84\%$) for myocardial ischemia [57, 58]. The primary reason for the underutilization of MCG seems to be purely economic: the total running costs of existing low-$T_c$ MCG systems are still too high for ordinary hospitals and clinics. This is true even though high quality MCGs can be generated in the absence of a bulky and expensive magnetically-shielded room [59]. The utilization of high-$T_c$ DC SQUID biomagnetometers for MCG leads to a significant reduction in running costs. Although they still require magnetic shielding, long-term savings make them a cost-effective alternative to low-$T_c$ MCG and perhaps even ECG.

Proof-of-principle MCG recordings with the two-channel high-$T_c$ SQUID system were performed in the MSR of the Franz-Volhard Hospital [53]. At frequencies below 10 Hz, an increase in environmental noise, which was associated with the reduced shielding factor of the MSR, was observed. In the 10–30 Hz frequency range, magnetic field signals caused by building vibrations were also observed. By using analog subtraction of the signal output of the top magnetometer (reference channel) from the signal output of the bottom magnetometer (signal channel), the parasitic low frequency noise determined by the low frequency signal, vibration effects, and 50 Hz interference was reduced by more than an order of magnitude, such that NEMI levels of below 80 fT Hz$^{-1}$ were reached at 1 Hz. The effective gradiometric resolution for the ‘electronic’ gradiometer was below 1 fT cm$^{-1}$ Hz$^{-1/2}$ at frequencies of above 100 Hz and below 2 fT cm$^{-1}$ Hz$^{-1/2}$ in the frequency range down to $\sim 4$ Hz. The white noise of the electronic gradiometer exceeded the noise of the individual magnetometers by a factor of approximately $\sqrt{2}$, in accordance with theoretical expectations.

MCG measurements were performed in the MSR with a volunteer having maximal peak-to-peak amplitude of $\sim 30$ pT for the magnetic signal of the heart. Both the magnetometer outputs and the electronic gradiometer signal were recorded simultaneously. The real-time result of subtraction of the reference channel is demonstrated in figure 7. A line frequency synchronous filter combined with a bandpass filter suppressed the residues of 50 Hz interference and its harmonics. The observed width of the noise trace of $\sim 500$ fT corresponds to an average NEMI of $\sim 16$ fT Hz$^{-1/2}$ in the measurement frequency bandwidth 0.03–125 Hz that covered the most significant frequency range of the heart signal.
recording time was 30 s per location.

Figure 7. Real-time MCG measured by the electronic gradiometer. The frequency bandwidth was 0.03–125 Hz. Reprinted from [34], with the permission of AIP Publishing.

Figure 8. Signal plots over the thorax show averaged MCG signals measured by the electronic gradiometer at 6 × 6 grid positions. The recording time was 30 s per location.

The system’s operating stability was demonstrated via measurements of MCG maps. Magnetic field measurements were carried out over a 6 × 6 square grid over the thorax with a distance of 4 cm between neighboring measurement positions in both directions (see figure 8). The total scan area was 20 cm × 20 cm. We used the simultaneously measured ECG channel as a trigger for averaging 30 s of MCG data at each measurement point. The low frequency (~0.1 Hz) drift of the magnetic field in the MSR and the effect of breathing caused some vertical offsets in the averaged differential signals, which can be seen in figure 7.

The total measurement time for all points was approximately 30 min. The magnetometers demonstrated stable operation and no change in noise performance over this time, allowing low noise 2D plots of the magnetic field distribution to be obtained. Equiinductional maps of magnetic field distribution were plotted on the basis of a two-dimensional interpolation at any moment of the heartbeat, as shown in figure 9. Interestingly, the magnitude of the MCG signal measured by the first order high-$T_c$ electronic gradiometer was less attenuated for measurement points towards the periphery of the scan area, as compared with corresponding data from a low-$T_c$ MCG system (based on second order gradiometer and Nb-wire pickup loops). This difference can be attributed to the increasing spatial filtering effect as the gradiometric order increases. The magnitude of the lead-field of a second order gradiometer decays more rapidly as a function of distance from the primary coil, when compared to that of a first order gradiometer with the same baseline.

3.2. Application of high-$T_c$ DC SQUID biomagnetometers for MEG

Brain disorders represent approximately 35% of human diseases and lead to enormous suffering for patients and their families world-wide. It was estimated that the total cost of brain disorders was about €800 billion in 2010 for Europe alone [60]. Understanding the origin of brain diseases and dysfunctions is complicated by its extreme complexity. The human brain contains approximately 100 billion neurons at maturity, all with relative arrangements, mutual connections and functions that change over time. A variety of neuroimaging modalities exists, such as functional magnetic resonance imaging and positron emission tomography for the study of hemodynamics and metabolism, respectively, with high spatial resolution. In order to explore and understand spatio-temporal brain dynamics, electroencephalography (EEG) and/or MEG are used to study neuronal activity. Both modalities provide direct access to electrophysiological activity with a superb temporal resolution of about one millisecond and are non-invasive. Today, they are employed for neurological and psychiatric research and provide valuable information about the spatio-temporal organization of the human brain. In contrast to EEG, MEG is known to provide better spatial resolution, as it does not depend on heterogeneity in conductivity across different head tissues (e.g., skull, brain, cerebrospinal fluid, etc.). However, when compared to MRI, the spatial resolution of MEG is worse and existing MEG technology needs further development to obtain reliable information from deeper brain areas with high spatial and temporal resolution. All commercial MEG systems are currently based on low-$T_c$ SQUIDs, which require liquid helium for cooling to operate at 4.2 K. In contrast to liquid nitrogen, the availability of helium cannot be guaranteed for the future. Moreover, the ever-increasing cost of liquid helium threatens wide acceptance of MEG as a diagnostic tool in clinical practice and results in pressure to develop new sensors for MEG research that would be independent of liquid helium. While cryocoolers placed next to a MEG system may reduce or even eliminate helium consumption significantly, such an approach has yet to reach maturity in terms of, e.g., routine clinical utilization, with a significant problem being additional noise contributions.

In recent years, improvements in high-$T_c$ SQUID technologies have shown that neuromagnetic signal recordings and results from post-processing obtained with a high-$T_c$ SQUID system are in good agreement with those obtained from a commercial low-$T_c$ MEG system [15, 20]. MEG measures magnetic field changes that originate from cortical
activity with field strengths in the range of a few hundred fT. A typical NEMI for low-$T_c$ MEG systems is about 4 fT Hz$^{-1/2}$, which is mainly limited by magnetic noise generated by thermal current fluctuations in the cryostat and the metal walls of the MSR and/or limited shielding of environmental noise by the MSR. For most routine MEG applications, a NEMI value of <10 fT Hz$^{-1/2}$ in the frequency range 1 Hz–1 kHz is sufficient and can be provided by high-$T_c$ SQUID biomagnetometers. These circumstances have justified our proof-of-principle MEG recordings using high-$T_c$ SQUID biomagnetometers, whose NEMI values are on that level [20, 54]. Importantly, our results indicate that the sensitivity of high-$T_c$ SQUIDs is ready for MEG source analysis [54].

In the study reported in [13] (see figure 10), MEG measurements of spontaneous brain activity were performed in the MSR formerly located at Imego AB in Gothenburg. Two of the ‘direct-coupled’ magnetometers were housed in separate cryostats. The head-to-sensor distance was estimated at ~1 mm for both channels. We studied the visual alpha (8–12 Hz) and sensorimotor alpha/mu (10–14 Hz) spontaneous brain rhythms via simultaneous single-shot recordings from O2 (visual/back right) and C4 (sensorimotor/right side) positions of the head with respect to the 10–20 coordinate system [61]. Both rhythms are well understood resting state neural oscillations, whose magnitudes are maximal when one is relaxed but alert. Visual alpha is strongest when the eyes are shut, while sensorimotor alpha/mu is strongest when there is a lack of both sensory input and motor control. Opening of the eyes suppresses visual alpha, whereas touch sensation and voluntary movement suppresses sensorimotor alpha/mu. Hand relaxation and flexure/closure were selected for modulating sensorimotor alpha/mu because the hand region occupies a large part of the sensory and motor cortices, thereby providing clear signals for recordings. The recording shown in figure 10 was 64 s long. During the first 32 s, the subject sat with open eyes and relaxed hands. After 32 s, the subject was verbally instructed to simultaneously close his eyes, flex his hands and hold that state for the remaining 32 s of the recording.

Time–frequency analysis (TFA) was applied to both raw biomagnetometer recordings. A boxcar window binned the data with a step size of 0.7 s and an epoch width of 3 s. Each epoch then underwent an FFT. Alpha/mu rhythms typically appear at 8–12 Hz, the specific peak frequency of which is subject-specific. Figure 10 shows spectrograms for the subject, whose alpha/mu rhythms peaked at around 11 Hz. As expected, the visual alpha wave signal recorded at O2/the back of the head is weak when the subject had open eyes and is strong upon closing them. The simultaneously-recorded C4/side of the head channel provides evidence of sensitivity.

Figure 9. Magnetic field distributions over the 20 cm $\times$ 20 cm scan area measured at moments of the (a) R-, (b) S- and (c) T-peaks of the MCG.

Figure 10. Time–frequency spectrograms of two-channel single-shot/un-averaged high-$T_c$ SQUID biomagnetometer recordings of the modulation of neural oscillations. Adapted from [13], with the permission of AIP Publishing. (a) The channel placed at O2/backside of the head records noise for the first 32 s of the recording, as the visual alpha signal is suppressed while the subject has open eyes. Upon closing of the eyes at 32 s and thereafter, clear alpha-band spontaneous brain rhythm activity is evident. (b) The channel placed at C4/side of the head (roughly over the sensorimotor cortex) records clear alpha/mu-band spontaneous brain rhythm activity as the subject sits with relaxed hands. Upon flexing/closing of the hands, this alpha/mu-band activity is suppressed below the noise floor of the recording.
to the sensorimotor alpha/mu rhythm, as the activity in that frequency range, while the subject is relaxed, is attenuated upon flexure/closer of the hands (also as expected). Despite the relatively high NEMI values of the simple single-layer ‘direct-coupled’ bicrystal-based JJ DC SQUID technology, this experiment demonstrated clear sensitivity to spontaneous brain activity with single-shot SNR levels of greater than 5.

In the studies reported in [20, 54], all MEG measurements were performed in the MSR of the Institute of Neurosciences and Medicine, Forschungszentrum Jülich, using both the high-\(T_c\) DC SQUID measurement system and the commercial 248 channel low-\(T_c\) MEG system ‘Magnes\(^\circ\) 3600 WH’ 4D-Neuroimaging. The analog output from the high-\(T_c\) system was connected to a 16-bit analog-to-digital converter port available on the low-\(T_c\) system with a sampling rate of 678 Hz and processed together with the signals from the low-\(T_c\) magnetometers. The NEMI of the high-\(T_c\) system was \(\sim 5 \text{fT Hz}^{-1/2}\) with the high-\(T_c\) magnetometer operating at 77 K in a cylindrical superconducting shield and \(\sim 7 \text{fT Hz}^{-1/2}\) (white noise level) at 77 K within the MSR. The low-\(T_c\) system had a NEMI of \(\sim 5 \text{fT Hz}^{-1/2}\) with the low-\(T_c\) SQUID operating at 4.2 K in the MSR. The head-to-sensor distances were about 15 mm and 19 mm for the high-\(T_c\) and low-\(T_c\) systems, respectively.

In 2014, the first successful source localization of neuromagnetic evoked responses was demonstrated using a single channel high-\(T_c\) SQUID [54]. In addition, TFA on resting state brain activity from visual areas revealed similar power spectral densities of spontaneous activity when compared to a commercial full-head low-\(T_c\) SQUID MEG system. A single subject (male, age 51, right handed) volunteered for both MEG experiments.

In order to investigate whether time–frequency-dependent oscillations recorded with a high-\(T_c\) SQUID system resemble those from a commercial low-\(T_c\) SQUID system, resting state brain activity was recorded in the same subjects with and without eyes closed using both high-\(T_c\) SQUID and low-\(T_c\) SQUID systems [54]. For the high-\(T_c\) SQUID measurements, spontaneous activity was recorded over the right occipital cortex. Neuromagnetic field changes were recorded continuously for 3 min with blocks of open/shut eyes for about 10 s, while the subject was asked (verbally) to open or close his eyes. The beginning of each block was documented by a change in the trigger code.

TFA was applied to spontaneous data recorded with the high-\(T_c\) and low-\(T_c\) SQUID systems using one channel located over the right occipital area of the subject. A total of nine epochs of 7 s duration were used to compute an average spectrogram, while all spectrograms were normalized to have unit variance prior to analysis for comparison between the low-\(T_c\) and high-\(T_c\) SQUID systems. In figure 11, clear changes in alpha power (8–12 Hz) are evident in both measurements using the low-\(T_c\) and high-\(T_c\) SQUID MEG systems. TFA of both data sets revealed spectrograms showing synchronization (increased power) and desynchronization (decreased power) of alpha rhythm upon closing and opening of the eyes, respectively, as expected.

For comparison between the low-\(T_c\) and high-\(T_c\) SQUID based power spectral densities, the MEG signals from both systems were normalized to unit variance prior to analysis. The peak-to-peak amplitude of the signal detected using the high-\(T_c\) system was slightly lower (\(\sim 500 \text{fT}\)) than the peak-to-peak amplitude of the signals shown in figure 10 due to the larger sensor-to-scalp standoff for the 16 mm flip-chip sensors. However, the SNR in the case of the flip-chip sensors is higher due to lower NEMI values of the sensors.

The experiments described above showed that the signal dynamics and power recorded using high-\(T_c\) SQUID sensors resemble those recorded using low-\(T_c\) MEG systems. However, in order to test and compare results between sensor technologies, the same brain signals need to be recorded from various positions around the source of interest (i.e., a given brain activation). This is trivial using conventional full-head MEG systems, whose sensors surround the head, but is a challenge when only one (or a few) high-\(T_c\) SQUID biomagnetometers are available. We developed a protocol for efficient benchmarking of single (or a few) channel magnetometers versus conventional MEG sensor technology. The protocol, which is detailed in [15], includes obtaining a recording using a standard MEG system, source localization based on that data, forward projection of the magnetic field generated by the source to the head surface, and recordings with the high-\(T_c\) SQUID biomagnetometer at head locations where the projected magnetic field magnitude is maximal. We also included EEG recordings (the magnitudes of which are maximal at positions on the head that are rotated by 90° with respect to the magnetic field maxima and therefore easily accessible) during all MEG sessions for assessing the stability of the activations.

We compared the low-\(T_c\) SQUIDs within the Elekta Neuromag TRIUX system (102 magnetometers with a 21 mm \(\times\) 21 mm pickup loop and 1–5 \text{fT Hz}^{-1/2} magnetic field resolution at 4.2 K and 10 Hz) to one of the ‘direct-coupled’ high-\(T_c\) SQUID biomagnetometers (9 mm \(\times\) 9 mm pickup loop and \(\sim 40 \text{fT Hz}^{-1/2}\) magnetic field noise at 77 K and 10 Hz). We selected the sensory N20m component induced by electrical stimulation of the median nerve (at the wrist) and the auditory P100m component induced by 1 kHz tones because both of these activations are known to be robust and to show little habituation. The recordings shown in figure 12 were performed inside the MSR at the Karolinska Institute NatMEG center in Stockholm. Low-\(T_c\) (blue) versus high-\(T_c\) (red) SQUID recordings were performed at the positions of maximal MEG signals. These positions are different for the low-\(T_c\) and high-\(T_c\) SQUID recordings because the sensor-to-scalp standoff is \(\sim 40 \text{mm}\) for the low-\(T_c\) system and \(\sim 5 \text{mm}\) for the high-\(T_c\) system. The difference between the positions of the sensors is the reason for the difference between the characteristic features of the MEG recordings. Figure 12 shows that for the auditory evoked fields the signal obtained using the high-\(T_c\) system is stronger than the auditory evoked field signal obtained using the high-\(T_c\) system by a factor 2.6–2.9. In the case of a somatosensory evoked field, the gain is between 3.4 and 4.2. Further experimental details can be found in [15].
A more comprehensive way to test and compare results obtained using high-$T_c$ SQUID biomagnetometers versus conventional low-$T_c$ MEG systems is to perform recordings from many head positions, such that source localization can be implemented using both sensor types. In [54], auditory evoked magnetic field responses were recorded in separate

Figure 11. Time frequency analysis of alpha-band spontaneous neural oscillations recorded above the right occipital area using a low-$T_c$ SQUID system (top row) and a high-$T_c$ SQUID system (bottom row) from one subject with eyes opened (left column) and closed (right column). For both systems, the spectrograms show enhanced alpha activity during time periods of eyes closed, while desynchronization of the alpha rhythm is evident during recordings with open eyes. For comparison between the low-$T_c$ and high-$T_c$ SQUID based power spectral densities, the MEG signals from both systems were normalized to unit variance prior to analysis. Reprinted from [55], with the permission of AIP Publishing.

Figure 12. Low-$T_c$ (blue) versus high-$T_c$ (red) SQUID recordings of the (a) auditory P100m and (b) somatosensory N20m activations. © 1969 IEEE. Adapted, with permission, from [15]. Blue bars indicated the expected field magnitude of the activations based on the expected gain in signal available when recording the neuromagnetic field at the scalp surface (i.e., with high-$T_c$ SQUIDs), in comparison to a few cm away (i.e., with low-$T_c$ SQUIDs). For the auditory evoked fields, the expected gain was a factor of 2.6–2.9. The magnitude of the high-$T_c$ recording is just below this range for the positive lobe of the field (pictured), and just above it for the negative lobe (not pictured). In the case of the somatosensory evoked field, the high-$T_c$ signal significantly exceeds the expected gain of 3.4–4.2 for both the negative (pictured) and the positive (not pictured) lobes.
runs and at 16 different locations above the left temporo-parietal area (the region surrounding the auditory cortex) using both the low-\(T_c\) and the high-\(T_c\) MEG system. These measurements were performed in Jülich using a single channel high-\(T_c\) MEG system equipped with 16 mm flip-chip high-\(T_c\) DC SQUID biomagnetometers and a 248-channel low-\(T_c\) MEG system.

The measurement locations for the high-\(T_c\) SQUID system were defined by the location of the electrode position of the EEG cap (EASYCAP GmbH, Berlin, Germany), which the subject was wearing without EEG electrodes (see figure 5(a)). In order to define the subject’s head, a coordinate system, including the coordinates of the electrode locations, the head surface and the electrode position of the EEG cap, were digitized using a 3D digitizer from Polhemus (Polhemus, 3Space/ Fastrak, USA). The cylindrical liquid nitrogen dewar from the high-\(T_c\) system was positioned above 16 different locations of the electrode mountings of the EEG cap. With the EEG cap placed between the head and the cryostat (see figure 5(a)), the distance between the subject’s head and the pickup coil was approximately 20 mm for all measurement locations.

During the experiments, 300 binaural stimulations were presented randomly (SOA \(6^{\pm}1.8\pm0.3\) s), while neuromagnetic brain responses were recorded. The intensity of the auditory click tones was adjusted to the comfortable hearing level of the subject. All MEG measurements were performed with the subject in the supine position. The neuromagnetic activity was continuously recorded with a sampling rate of 678.17 Hz and a bandwidth ranging from DC to 200 Hz. All tone onsets were shifted by −19 ms due to the delay of the sound delivery system, while an additional delay of 6 ms was subtracted from the high-\(T_c\) SQUID measurement due to the constant delay in the high-\(T_c\) SQUID readout electronics. For comparison, responses to auditory stimulation from the entire brain were acquired using a whole-head MEG system equipped with 248 low-\(T_c\) SQUID magnetometers.

A specific analysis procedure was implemented with respect to source localization of the auditory-evoked field recorded by the high-\(T_c\) SQUID system. The identification of unusually large amplitudes (e.g., due to artifacts) in the signal of the high-\(T_c\) and low-\(T_c\) SQUID recordings was performed on the raw data by using a peak-to-peak measure applied to the full data set with a sliding window of length 200 ms. A bandpass filter ranging from 3 to 30 Hz as well as a notch filter at the power line frequency of 50 Hz and harmonics were applied. Trials around the stimulus onset were defined with 1 s post- and 400 ms pre-stimulus. The number of trials included for further analysis after discarding those with artifacts or unusually large amplitudes was, on average, 293 and 294 for the low-\(T_c\) and high-\(T_c\) SQUID recordings, respectively. Finally, all trials were averaged based on stimulus onset time. For comparison of the results from source analysis between the two systems, positions from the high-\(T_c\) SQUID measurements were aligned to low-\(T_c\) SQUID sensors that best matched the location of the high-\(T_c\) SQUID recording.

A dipolar field pattern can be seen in both the low-\(T_c\) SQUID and the high-\(T_c\) SQUID topographic field maps (see insets over the head sketches in figure 13). Isocurves of the evoked fields are shown for a peak latency of about 87 ms after the stimulus onset. The field distribution measured using the high-\(T_c\) SQUID system at 16 different positions resembles the field map of the low-\(T_c\) SQUID system in this area and clearly covers the zero-field-line that roughly indicates the location of the source. The left part of figure 13 shows MEG signal averages from the left hemisphere in response to auditory stimulation. The top panel shows a signal average using 50 channels over the left temporal cortex, as obtained by the whole head low-\(T_c\) SQUID magnetometer system (blue). The bottom panel shows signals from the same subject obtained using a single high-\(T_c\) SQUID at 16 different positions (red). The time range highlighted in gray was used for source analysis. The traces obtained from both the high-\(T_c\) and the low-\(T_c\) systems demonstrate similar SNR, despite the 18% difference in the operating temperatures of the sensors.

The single dipole fit algorithm, as provided by the 4D-Neuroimaging MEG system, was applied to localize the sources of the dipolar fields that were recorded \(^{62}\). Analysis of the high-\(T_c\) SQUID measurements included MEG signals from the 16 different positions (figure 13). For analysis of the low-\(T_c\) SQUID recordings, the system pre-defined group of 50 channels covering the left temporoparietal area was used. The larger number of channels was kept for source analysis of the low-\(T_c\) system data because it serves as a reference.

Estimation of the dipolar source was performed using a time window of 50 ms between 60 and 110 ms. The dipole with the best goodness-of-fit value from the low-\(T_c\) SQUID measurement was chosen for comparison and was found to be maximal 104 ms after stimulus onset. Analysis of the high-\(T_c\) and low-\(T_c\) SQUID recordings revealed a dipolar source with the same latency as expected (figure 14). The location of this source, as determined using the single high-\(T_c\) SQUID system, is in very good agreement to that determined using the low-\(T_c\) system. Both the location and the direction of the source estimations are comparable. The high-\(T_c\) SQUID source analysis revealed a displacement of only 7 mm compared to the results obtained using the low-\(T_c\) SQUID system \(^{54}\).

A possible further development of the high-\(T_c\) MEG system is to use several ten-channel systems with individual cryostats. Each cryostat can then enclose, for example, seven signal channels and three reference channels. A photograph of such a cryostat and a sketch of a ten-channel MEG system that is under construction are shown in figures 15(a) and (b), respectively. The \(\sim 11\) l commercial cryostat from Tristan Technologies Inc. (originally intended for liquid helium) holds liquid nitrogen for more than two weeks with an evaporation rate of approximately 0.8 \(1d^{-1}\). A liquid nitrogen evaporation rate of below 0.3 \(1d^{-1}\) is expected for a cryostat of similar size that is specifically designed to hold liquid nitrogen. Such a seven-signal-channel system configuration will be used for the investigation of a region of human brain that is related to a specific functional activity.

\(^{6}\) Stimulus onset asynchrony (SOA) describes the average time from one stimulus to the other, whereas onset times are randomized.
Figure 13. Comparison of auditory evoked MEG averaged signals acquired at a similar location over the left hemisphere using low-\(T_c\) (50 positions, blue, upper curve) and high-\(T_c\) (16 positions, red, lower curves) SQUID systems. The time range highlighted in gray was used for source analysis. Adapted from [55], with the permission of AIP Publishing.

Figure 14. Results from source analysis of auditory evoked neuromagnetic field responses, as recorded using high-\(T_c\) SQUID (red circle) and low-\(T_c\) SQUID (blue triangle) systems. The location of the dipolar source from the high-\(T_c\) SQUID system (red) is, as expected, in the region of the left primary auditory cortex and is in very good agreement with source analysis from data recorded using the low-\(T_c\) SQUID system (blue). Middle image reprinted from [55], with the permission of AIP Publishing.

Figure 15. (a) Photograph of a cryostat intended for a ten-channel MEG system fixed on an adjustable cryostat holder in a magnetically-shielded room in the Peter Grünberg Institute, Forschungszentrum Jülich GmbH. (b) Sketch of a ten-channel MEG system that is under construction.
3.3. High-\(T_c\) DC SQUID biomagnetometers and MNPs

Ferrimagnetic crystals of magnetite (Fe\(_3\)O\(_4\)) that are below 100 nm in size occur naturally in bacteria, protists, animals and humans due to biomineralization (see [1, 63, 64] and references therein). High-\(T_c\) DC SQUIDs can easily resolve the magnetic fields produced by the motion of even a single living magnetotactic bacterium [65]. Functionalized MNPs of magnetite are used for the magnetic hyperthermia treatment of cancer and serve for targeted drug delivery to increase drug concentration in tissues or because they can be transported through the vascular system and concentrated at a particular point of the body with the aid of a locally applied magnetic field [1, 66–68]. Antibody-conjugated MNPs are able to find (in vivo and/or in vitro) proteins that are specific for cancer, Alzheimer, and Parkinson diseases, to serve as their magnetic biomarker, and to diagnose these diseases at an early stage with the help of high-\(T_c\) SQUIDs [69–71].

Chalmers has developed a one-step biomolecular assay that is based on sensitivity to changes in fluid-suspended MNP Brownian relaxation dynamics in the frequency and time domains [55]. For this experiment, biofunctionalized multi-core particles containing single domains of CoFe\(_2\)O\(_4\) (Chemicell GmbH) with a median hydrodynamic radius of 50 nm were used. The MNPs were functionalized with streptavidin with biotinylated monoclonal antibodies. A prostate-specific antigen (PSA) assay was targeted as a model system. MNPs can bind 80 pmol of biotinylated protein per mg of MNPs, which corresponds to approximately 30 PSA10 molecules per MNP. PSA10 is a monoclonal antibody for PSA and acts as a capturing antibody for the specific targeting of PSA. In this study, a ‘direct-coupled’ high-\(T_c\) DC SQUID gradiometer with balance 93, spectral flux noise 4.5 \(\mu\)Phi/Hz\(^{-1/2}\) at 1 kHz, and a gradient resolution of \(\sim 2.1\) pT cm\(^{-1}\) Hz\(^{-1/2}\) was used [26]. The same ILK Dresden cryostat was used as in the Chalmers ‘direct-coupled’ MEG experiments. For such measurements of fluid-suspended MNPs, proximity between the sensor and the room temperature environment is particularly powerful. The size of the gradiometer (baseline 3 mm) was small enough to be used effectively without magnetic shielding, while having sensitivity to extremely small volumes (as little as 1 \(\mu\)l) of fluid-suspended MNPs. Despite the size of this volume, the roughly \(1 \times 10^8\) MNPs have a total surface area of several square centimetres. This is significantly larger than standard well-based biomolecular assays, whose binding areas (i.e., ‘sensing areas’) are typically no larger than 1 cm\(^2\). The estimated MNP content sensitivity of \(\sim 50\) ng Hz\(^{-1/2}\) (\(\sim 1.1 \times 10^8\) MNPs) was limited by the flux sensitivity of the gradiometer of \(\sim 100\) \(\mu\)Phi Hz\(^{-1/2}\) at 2 Hz. Figure 16 shows the sensitivity of the measurements to varying analyte concentrations as surface binding events slow the MNP relaxation. The samples were measured in the frequency domain (figure 16(a)) using AC-susceptometry at different frequencies (frequency dependence of the complex susceptibility) and in the time domain (figure 16(b)) using magnetorelaxometry and a pulsed magnetic field. As the concentration of analyte increases, the average Brownian relaxation slows, shifting the peak of the imaginary part of the complex susceptibility (figure 16(a)) down in frequency and slowing the relaxation dynamics in the time domain (figure 16(b)).

The minimum volume of the liquid containing the MNPs required to detect a signal was about 0.3 nl at a particle number density of \(1.8 \times 10^{15}\) g\(^{-1}\). This volume corresponds to \(2.2 \times 10^7\) MNPs with a total magnetic moment of 0.3 pA m\(^2\) and an average magnetic dipole moment per particle of \(4.8 \times 10^{-22}\) A m\(^2\). The observed remnant signal was probably due to alignment of the magnetic moments of the MNPs in the Earth’s magnetic field. The estimated ultimate biomolecular sensitivity for this measurement system is \(\sim 8 \times 10^8\) molecules in a 2 \(\mu\)l sample volume, corresponding to a density of MNPs of \(\sim 100\) ng ml\(^{-1}\). This measurement system can be
adapted to other disease markers, for which trapping antibodies are available.

4. Discussion

Despite the discovery of high temperature superconductors by Bednorz and Müller in 1986 [72] and subsequent developments of high-$T_c$ SQUIDs that are capable of operation at liquid nitrogen temperature, all commercial MCG and MEG devices are still presently based on low-$T_c$ SQUIDs that require liquid helium for operation at 4.2 K. Nevertheless, sensors that function at liquid nitrogen temperature (77 K) have significant economic and technical advantages. Both earlier and more recent publications have reported successful recordings of MCG and MEG that utilize single-channel or multi-channel systems [73–78] based on direct-coupled high-$T_c$ SQUID magnetometers with bicrystal JJs. MCG systems that are based on direct-coupled high-$T_c$ SQUIDs demonstrate sufficiently high SNR in MSRs because typical MCG signals are relatively large. Outside MSRs, the operation of high-$T_c$ SQUID magnetometers is severely degraded due to electromagnetic interference, which leads to instability and too low SNR of MCG signals in most locations. Electromagnetic interference can be avoided in the case of well-balanced second order gradiometric low-$T_c$ MCG systems that are able to operate well outside MSRs [59]. Direct-coupled high-$T_c$ SQUID magnetometers also have advantages for applications to MNPs in bioassays and magnetic label counting electronics is used, as well as relatively small crosstalk values. Crosstalk between sensors is critical for accurate representation of the spatial distribution of the magnetic field in both MCG and MEG. The magnetic field of the feedback coil decays as a function of its area $\sim A^{-3/2}$. The feedback signal for SQUIDs can be injected directly into the A $\sim 0.01$ mm$^2$ area of the SQUID loop. In the case of atomic magnetometers, the feedback coil should provide a homogeneous field over a cross-sectional area of $\sim 300$ mm$^2$ [9]. As such, the radius of equivalent crosstalk (i.e., the distance between sensors with equal crosstalk) for atomic magnetometer sensors can be as much as $\sim 5 \times 10^6$ larger than for SQUIDs. While all SQUIDs require low operating temperatures, high-$T_c$ SQUIDs are far less limited by this restriction when compared to low-$T_c$ SQUIDs.

Crosstalk and cooling aside, the primary advantage of using high-$T_c$ SQUIDs for MEG and MCG is that long-standing experience in SQUID sensor, readout electronics, and dewar technology exists. Therefore, a potential shift from low-$T_c$ to high-$T_c$ SQUID technology is expected to be straightforward. Theoretically, the field resolution of high-$T_c$ SQUID magnetometers can be improved further. A magnetic field resolution of well below 1 fT Hz$^{-1/2}$ at 77 K should be achievable by using electron beam lithography to pattern JJs, a dual SQUID layout, flux transformers with larger pickup loops, and liquid nitrogen pumped down to its triple point temperature of 63.15 K. There is also the possibility of extending the current technology further for the preparation of multilayer high-$T_c$ DC SQUID planar gradiometers of first or second order on single crystal MgO substrates that are up to $\sim 10$ cm long. Such a development can be useful for MEG recordings where the magnetically shielded environment is relatively weak. High-$T_c$ SQUIDs with novel graphoepitaxial step-edge junctions can be fabricated at lower cost and with improved reproducibility, when compared to the previously widely used bicrystal technology. Perhaps more importantly in terms of biomagnetometry, high-$T_c$ SQUIDs also enable a significant reduction in distance between the pickup coil and the subject’s head (from $\sim 20$ to $\sim 3$ mm or less) or chest, leading to improved MEG/MCG recordings when compared to the state-of-the-art.

5. Summary

In summary, high-$T_c$ SQUID biomagnetometers have been shown to demonstrate their usefulness for the two most important biomedical applications: MCG and MEG. The results encourage and pave the way for the development of multichannel high-$T_c$ SQUID based MCG and MEG systems. In the long term, we expect low-$T_c$ DC SQUIDs to be replaced by graphoepitaxial step-edge-junction-based high-$T_c$ technology for many applications, including future biomagnetic systems. Such an upgrade would make future multichannel systems independent of helium supplies, more user-friendly, and perhaps even more sensitive than what is available today. For example, we expect that a helium-free MEG system may save up to about (a few MHz [11]) and large dynamic range (>100 dB) if flux counting electronics is used, as well as relatively small crosstalk values.
100,000 \text{\euro\ yr}^{-1} due to a reduction in the operating cost for each system. More importantly, the availability of liquid nitrogen is guaranteed, in contrast to liquid helium. High-$T_c$ SQUIDs with novel graphoepitaxial step-edge junctions \cite{20, 22, 29, 31} promise better superconducting parameters, better reproducibility and lower cost when compared to previously widely used high-$T_c$ SQUIDs with less available bicrystal JJs. When compared to other sensor technologies, the advantage of using SQUIDs is their combination of high sensitivity (a few fT Hz$^{-1/2}$), frequency bandwidth (a few MHz \cite{11}) and dynamic range (>100 dB), as well as relatively small crosstalk values.

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