In utero Diagnosis of Long QT Syndrome: Challenges, Progress, and the Future

Suhong Yu1 and Ronald Wakai2

1Department of Radiation Oncology, University of Rochester, New York, USA
2Department of Medical Physics, University of Wisconsin-Madison, WI, USA

Corresponding author: Suhong Yu, Department of Radiation Oncology, University of Rochester, New York, Tel: +1 585-275-2121; E-mail: yusuhong@gmail.com

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Introduction

Congenital long QT syndrome (LQTS) is an inherited ion channel disorder caused by gene mutations that encode for cardiac ion channels. The incidence of inherited LQTS in the general population is estimated to be 1:2000 (0.05%) [1]. LQTS manifests as prolonged corrected QT interval (QTc) on the electrocardiogram (ECG) and magnetocardiogram (MCG), and it is strongly associated with cardiomyopathies, channelopathies, and sudden death at all ages [2-5]. Sudden death results from susceptibility to Torsade de Pointes (TdP), a highly lethal ventricular tachycardia. Little is known about LQTS in utero, these capabilities can be lifesaving. We foresee a promising future for in utero diagnosis of LQTS and other life-threatening fetal rhythm disorders.

Challenges and Progress

Assessment of repolarization using MCG is considerably more difficult in the fetus than in the neonate or adult due to the lower amplitude of the signal and the presence of large interference, primarily from the maternal MCG. Signal processing techniques are required to remove the interference and facilitate detection of abnormalities such as QTc prolongation and T-wave alternans (TWA), i.e. beat-to-beat variation in the amplitude of the T-wave [17-19]. TWA is a rare, but significant, rhythm pattern indicative of cardiac instability.

In a recent fetal MCG study, the electrophysiology of LQTS in utero was characterized for the first time in a sizeable cohort, consisting of 30 fetuses at risk of LQTS [20]. Heart rate, waveform intervals, T-wave morphology, initiation /termination patterns of TdP, and TWA were assessed. Fetal MCG demonstrated high diagnostic and prognostic value. Based on assessment of QTc interval (QTc > 490 ms), fetal MCG was able to identify the fetuses that tested positive for LQTS with high accuracy (89%). Low-for-gestational age heart rate (< 3%) was also associated with fetal LQTS. Some fetuses diagnosed with LQTS had only low fetal heart rate and no family history of LQTS at the time of referral. In several such cases, LQTS was subsequently found in 1st-degree relatives who underwent testing as a result of the fetal MCG diagnosis. The fetal MCG findings also showed high prognostic value. Subjects that had TdP as fetuses or newborns showed the longest values of QTc (>600ms). TdP was also associated with other rare findings, including 2nd-degree AV block, TWA, and QRS alternans. Lastly, definitive detection of TdP was critical for guidance of in utero therapy, consisting of administration of magnesium and lidocaine, which was highly effective at controlling or abolishing TdP.

The Future

While the efficacy of fetal MCG has become more widely recognized, the high cost of the technology, which is based on superconducting sensors known as SQUID magnetometers, has limited its widespread use. This situation will likely change in the near future due to a recent breakthrough in atomic magnetometry, leading to the development of the so-called SERF (spin exchange relaxation free) magnetometer [21,22]. The SERF magnetometer is the first alternative device with sensitivity equal to or better than that of a SQUID magnetometer. Its main advantage, however, is low cost. The required optical components are inexpensive because they are already used in commercial products, e.g. DVD players; thus, atomic magnetometers can reduce the cost of fetal MCG detectors by nearly an order of magnitude. Promising results have already been published, and commercial systems based on atomic magnetometers will be realized in the near future [23].

Fetal MCG is an enabling technology for the in utero detection and management of LQTS. Due to the ability to effectively treat TdP in utero, these capabilities can be lifesaving. We foresee a promising future for in utero diagnosis of LQTS and other life-threatening fetal rhythm disorders.

References


