

Low Power Multiparameter Biopotential Amplifier System

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Abstract: Development of in vivo electronic devices has been the top priority of medical device development for the past few years. In the context, diagnosis of the bio potential system of human body before and after the treatment plays a lead role. This paper is about a low power multiparameter multichannel biopotential system which has its primary application in as front end circuitry for patient monitoring systems such as polysomnography, ICU monitoring etc. the system has its key features as low power operation, durability, battery based operation, high noise rejection capabilities, support for wireless integration, small size etc. The described system is an all in kind of system for acquiring and measurement of biopotential signals like ECG, EMG, EEG, EOG etc which are essentials of a PSG system.

Keywords: Diagnosis, biopotential system, data acquisition, wireless integration

1. Introduction

Advances in low power microelectronic devices over the years have given a rapid boost in the field of biomedical instrumentation. Bioelectronics interfaces that are miniaturized, light weight, low power, compatible with minimum interface requirements etc has been come out research and development laboratories in the mean time. Nowadays medical device development is undergoing dramatic changes absorbing advancements in research areas such as wireless automation, transcutaneous energy transfer, on invasive cardiac support devices delivery, total artificial heart etc.

Conventional block diagram of a biopotential recording system is shown in figure 1. For decades monolithic amplifiers were used for measuring and recording electrophysiological signals. Due to the large time constant that was inherently present in the monolithic amplifier's dynamics prevent the sharing of a single electrode between multiple electrodes. So multiple amplifiers, one per channel is used typically for multichannel systems.

Figure 1 explains the basic block diagram of a data acquisition front end. The system consists of a Data Acquisition System coupled with the electrodes to pick up vital signals from the human body. The system is equipped with an analog front end which facilitates the signal extraction and conditioning.

The system contains bioelectrodes depending upon the type of biopotential that is to be acquired, front end instrumentation amplifiers for the initial pick up of the signals, a signal conditioning circuit for noise removal, gain etc. The signal from the signal conditioning circuit is converted to a digital signal by means of an analog to digital converter (ADC). Since we need to transmit the signal via any serial communication means we can opt for an ADC inside any of the microcontroller since it has dedicated serial communication capabilities.

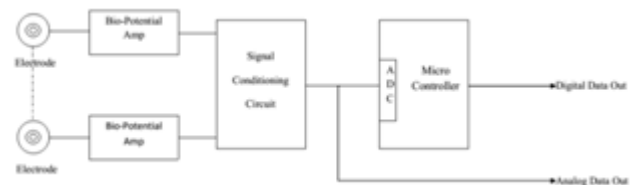


Figure 1: Biopotential Acquiring System

Table 1 shows the electrical characteristics of commonly acquired electrophysiological signals that are commonly acquired during diagnosis and treatment. Single unit recording, provides finest spatial resolution of the signals but they typically incur relatively higher power consumption due to wide amplifier bandwidth required and high resulting data rate.

Table 1: Characteristics of Electrophysiological Signals

	Bandwidth	Amplitude	Invasiveness
Single Unit			
LFP (Local field Potential)	<200 Hz	<5mV	Moderate
EEG(Electroencephalography)	<100 Hz	10-20µV	Non Invasive
ECG(Electrocardiography)	0.5Hz-120Hz	0.1-5mV	Non Invasive
EMG(Electromyography)	0-500 Hz	0-10mV	Both Ways

Technical specifications of the biopotential signals are to be considered while designing amplitude based recording amplifier. Cellular action potentials which fall in frequency range of 100 Hz to 7 KHz band have amplitude of up to 500 µv. Local Field Signals that come under low frequency band (<1 Hz) typically have an amplitude of up to 5 mV. There are some considerations in analog signal processing while discussing about bio potential amplifiers. Low amplitude levels of biopotential signals limit the gain of biosignal amplifiers to around 100 up to a few KHz. DC offsets which are introduced by artifacts at electrode tissue interface implies a need for offset cancellation or ac coupling. To limit the signal attenuation at electrode tissue interface the input impedance of the amplifiers should be very high in order of few MΩs. to contain the rise in temperature due to power dissipation the amplifier should draw only a minimal current. To deal with interference and power supply noise which are inevitable in amplifier designs sufficient Common Mode Rejection and Power supply rejection has to be ensured.

2. Bio Potential amplifiers

A biopotential amplifier's main function is to pick any signal of biological origin and amplify it so that it can be further processed analyzed, displayed or recorded. Normally those amplifiers alter the amplitude levels and there by power also so these can be called as voltage amplifiers or power amplifiers as well. Biopotential amplifiers when used for isolating the load, they only provide current gain leaving the voltage levels intact.

Most often bipolar electrodes are used to acquire biopotential signals. Speaking with respect to ground these electrodes will be electrically located in a symmetric manner. That makes the choice of amplifier to be a differential one. Distortions in the symmetry with respect to ground can introduce common mode voltage fluctuations in the amplifier so the amplifier should have high common mode rejection ratios to deal with the interference due to common mode signal. Biopotential amplifiers have additional requirements that are application specific to various signal types and its characteristics which are discussed below.

2.1 ECG-Electrocardiograph Amplifier

Heart health of a person can be easily identified from ECG waveform like arterial fibrillation can be detected from distortions in P wave.

2.1.1 Basic Principles of ECG

Small electrical changes are introduced in human skin when the heart muscle depolarizes during the heart beat which is detected and amplified by the ECG amplifier. There is a membrane potential across the cell membrane usually called as cell potential which is usually negative when the heart is at rest. Through influx of positive ions Na^+ and Ca^{++} will decrease the negative charge towards zero which activates the mechanisms in the cell that causes cell to contract is called depolarization. During each heartbeat a consistently paced wave of depolarization which is generated by the SA node-sinoartial node spreads throughout the atrium through purkinjee fibres to the ventricles. This is reflected as small rises and falls in the potential between electrodes placed across heart .this is displayed as a wavy line either on a screen or on a paper.[1][2]

The basic ECG signal will look like in Fig :2

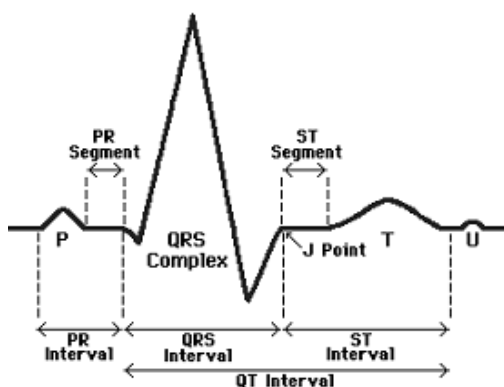


Figure 2: ECG Signal

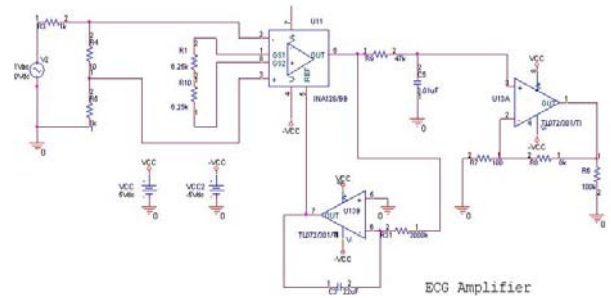


Figure 3: Schematic Diagram of the ECG amplifier

2.1.2 EMG- Electromyograph Amplifier

Electromyography (EMG) measures muscle response or electrical activity in response to a nerve's stimulation of the muscle. The test is used to help detect neuromuscular abnormalities. Instead of conventional needle electrodes we are using surface electrodes along with the designed highly versatile bio amplifier system to pick up the surface EMG signals. The electrical activity picked up by the electrodes is then displayed on an oscilloscope (a monitor that displays electrical activity in the form of waves). An audio-amplifier is used so the activity can be heard. EMG measures the electrical activity of muscle during rest, slight contraction and forceful contraction. Muscle tissue does not normally produce electrical signals during rest.[3]

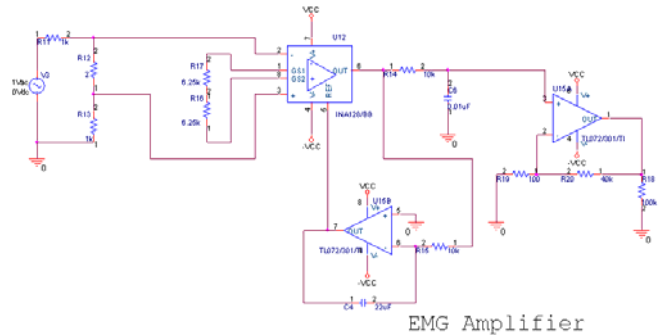
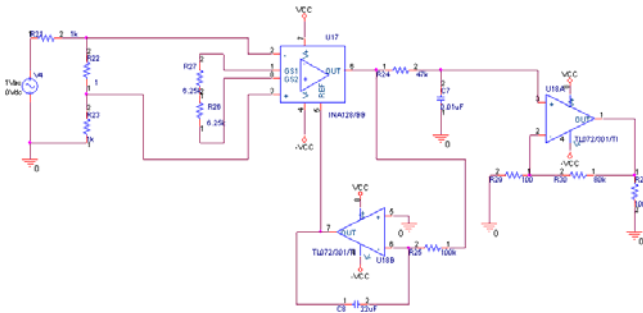


Figure 4: Schematic Diagram of the EMG amplifier.

2.2.3 EEG –Electroencephalograph

Electroencephalography (EEG) is the recording brains electrical activity over the scalp. Neurons in the brain produce voltage level variation due to ionic flow variations and thus producing EEG. In clinical contexts, EEG refers to the recording of the brain's spontaneous electrical activity over a short period of time, usually 20–40 minutes, as recorded from multiple electrodes placed on the scalp. Generally the spectral content in the EEG-ie the type of neural oscillations inside brain is taken into account for diagnosis. In neurology, the studies in EEG are mainly on Epileptic studies since variations in EEG signals are clearly depicted in case of Epilepsy. EEG can be used as a secondary diagnosis method for, coma, encephalopathies, and brain death. For studies related to Sleep disorders and sleep apnea EEG signals are being put on to use widely. The detection of sleep and differentiation between the REM and TEM stages of sleep is identified from the alpha waves of EEG only. As a first line method EEG can be used as a tagline feature for detection of tumors, stroke and other focal; brain disorders. Despite limited spatial resolution, EEG continues to be a valuable tool for research and diagnosis, especially when

millisecond-range temporal resolution (not possible with CT or MRI) is required.[2][3]



EEG Amplifier

Figure 5: Schematic Diagram of EEG amplifier

3. Simulation Results

The simulation results of three amplifiers are done using Pspice tool from cadence. The transient response and ac sweep analysis were performed to verify the amplification factor and frequency responses which are shown in the figures below.

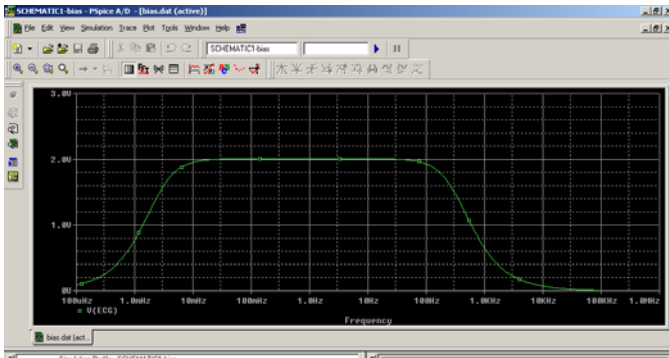


Figure 6: Frequency Response Curve of ECG amplifier

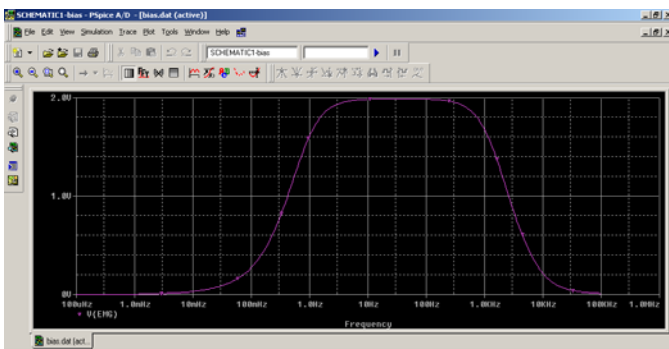


Figure 7: Frequency Response Curve of EMG Signal

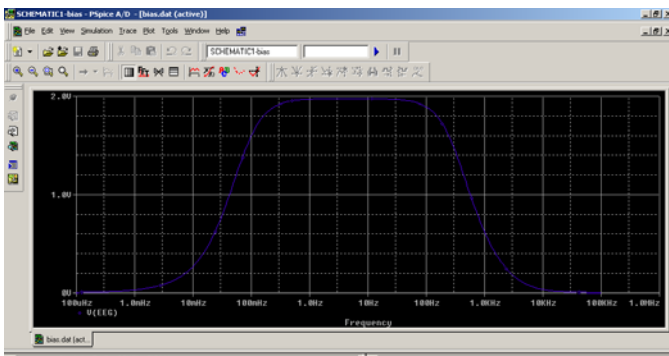


Figure 8: Frequency Response Curve of EEG Signal

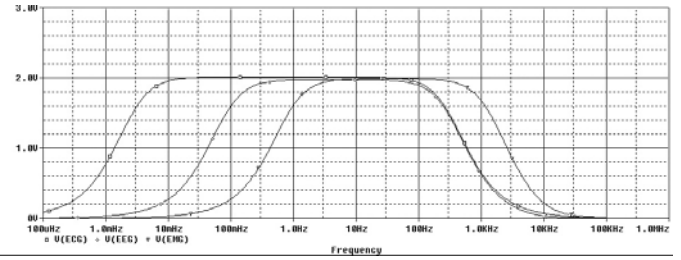


Figure 9: Comparison between frequency response curve of ECG, EEG and EMG signals.

4. PCB Design

The PCB of Bio-signal Amplifier is Double Sided one with dimension 15 x 18 x 1.6 mm. The PCB is provided with a mounting hole at the centre, so that PCB can use in a stacked manner for multiple channel application. The material used is FR4 with 35 uM copper thicknesses. The surface finish used is HASL (Sn-Pb) (Hot Air Solder Leveling). The PCB is routed with trace width 8 mil for signals and 12 mil for Power with a Track to Track spacing of 8 mil. The VIA used is 12/24 (drill/pad size, in mils). Thermal relief spoke width is 12 mil. The ground is poured on both side of the board for better connectivity and noise immunity. [8]

The PCB is designed in such way that to meet signal integrity needs. The board contains two connectors input and output connector and the power connector. The input lines are routed differentially for better noise immunity. An additional provision for shielding the circuit is also provided on the board through the mounting hole. So we can easily connect the PCB to chassis using a M2 earthing tag.

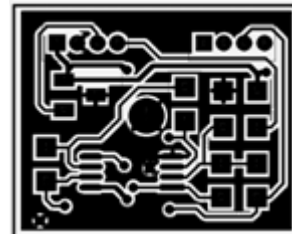


Figure 10: Bottom Layer Routing of PCB

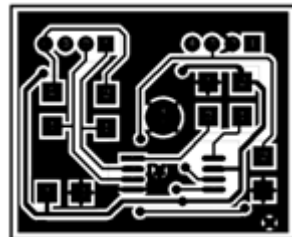


Figure 11: Top Layer Routing of PCB

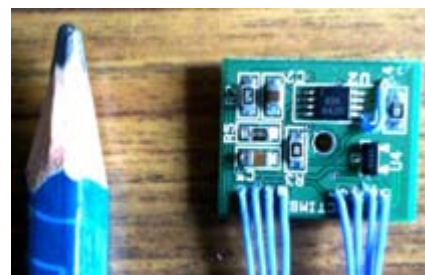


Figure 12: Actual Populated PCB. (Size comparison with a standard pencil tip)

5. Results

The PCB is fabricated as a four layer PCB with two routing layers and two power planes. The circuit was tested clinically with volunteers and indigenously developed electrodes that are developed in our laboratory only. Fig 13 shows the actual screen shot of an ECG wave form on Tektronix DSO.

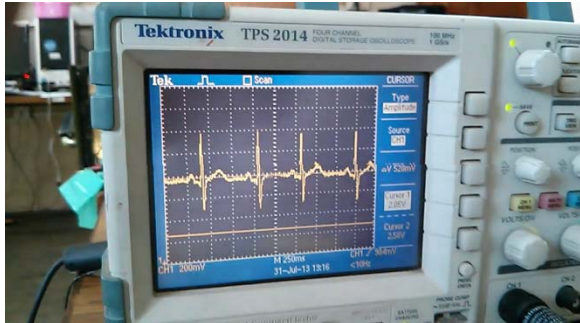


Figure 13: Real time acquisition on DSO.

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Author Profile



Jithin Krishnan received his M Tech Degree from Cochin University of Science and Technology in 2012. His area of interests is VLSI, Embedded systems and Bio Medical Instrumentation. He is currently doing research in biomedical instrumentation focusing on data acquisition systems and sophisticated medical devices development. He is a IEEE member for more than 5 years and is currently review committee member for IEEE Standards Association for Asia-Pacific.



Dr. Niranjana D. Khambete has B. E. in Instrumentation Engineering, M. Tech. in Biomedical Engineering and PhD in Engineering and has been actively contributing to the Medical Device R&D needs of the country for last two decades. He worked as a team leader to successfully complete the indigenous technology development and commercialization of Concentric Needle Electrodes. On research front, Dr. Khambete works on movement artifact free apnea monitors for early detection of sleep disordered breathing and bio-impedance spectroscopic measurement systems for early detection of cervical cancer in women and monitoring cell growth in tissue engineered organs. In recent years, Dr. Khambete has also focused his efforts towards spreading awareness about the Clinical Engineering Profession and promoting safe use of medical equipment in hospitals. In 2010, he received the WHO Patient Safety Award for his paper on medical device safety in hospitals at a conference in London. In 2011, the American College of Clinical Engineering gave him Antonio Hernandez International Clinical Engineering Award, for his leadership and commitment to the growth of Clinical Engineering Profession in India.



Biju Benjamin is working as Technical assistant (Instruments) in Biomedical Technology Wing Sreechitra Tirunal Institute for Medical Sciences & Technology, Thiruvananthapuram, Kerala. For Last 8 years he has been working in the Field of Electronics Testing, Simulation & PCB Design.



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