sEMG Interface Design for Locomotion Identification

Rohit Gupta, Ravinder Agarwal

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Abstract—Surface electromyographic (sEMG) signal has the potential to identify the human activities and intention. This potential is further exploited to control the artificial limbs using the sEMG signal from residual limbs of amputees. The paper deals with the development of multichannel cost efficient sEMG signal interface for research application, along with evaluation of proposed class dependent statistical approach of the feature selection method. The sEMG signal acquisition interface was developed using ADS1298 of Texas Instruments, which is a front-end interface integrated circuit for ECG application. Further, the sEMG signal is recorded from two lower limb muscles for three locomotions namely: Plane Walk (PW), Stair Ascending (SA), Stair Descending (SD). A class dependent statistical approach is proposed for feature selection and also its performance is compared with 12 preexisting feature vectors. To make the study more extensive, performance of five different types of classifiers are compared. The outcome of the current piece of work proves the suitability of the proposed feature selection algorithm for locomotion recognition, as compared to other existing feature vectors. The SVM Classifier is found as the outperformed classifier among compared classifiers with an average recognition accuracy of 97.40%. Feature vector selection emerges as the most dominant factor affecting the classification performance as it holds 51.51% of the total variance in classification accuracy. The results demonstrate the potentials of the developed sEMG signal acquisition interface along with the proposed feature selection algorithm.

Keywords—Classifiers, feature selection, locomotion, sEMG.

I. INTRODUCTION

CURFACE electromyography (sEMG) signals have a Dparamount importance and potential in the field of robotics, it is widely applied to control assistive and devices like prosthetics, rehabilitative exoskeletons, wheelchairs etc. The controlling of such devices is done by identifying the insight information in the signal [1]. To exploit the above-mentioned property of the sEMG signal many industries developed sophisticated and high cost sEMG measuring instruments like Nexus 10, Trigno etc. Many research groups have developed their own sEMG interface systems for laboratory use [2]-[4]. Earlier developed systems exploited instrumentation amplifiers e.g. AMP04FPZ, INA114, INA128P as a component [4]-[6], but the major disadvantage is that, for each channel one dedicated IC is used, which increase the component bourdon on developed instrument. Some of the researchers proposed active electrode system and systems with lesser circuitry [7], [8] but in these

systems dedicated cables and interface increase the cost of the system.

The hidden information in the sEMG signal is treated as feature of a particular signal, so that after signal acquisition, the appropriate signal processing techniques need to be applied, a trade off should be made between information loss and excess/vague information retention. Loss in information carried by the signal degrade the quality of the signal, such as the identification of functionality in the signal; whereas the excess information acts as noise that will ultimately retrench the performance of the system [9]. In the recent past, various researchers have applied different combinations of extracted features and pattern recognition algorithms and postprocessing techniques for the application of activity recognition of both upper and lower limbs [1], [3], [10]-[12]. The features utilized by previous researchers are mainly from the time domain, frequency domain and time-frequency domain (wavelet transform). Out of these three domains, the time domain based features gained popularity, as these are computationally inexpensive, easy to implement and provide better results [13]–[17]. Multiple studies had been performed by various researchers on activity identification using sEMG signal or mechanical sensors or both, and the majority of them are on upper limb activity identification [18], but in past couple of decades, recognizable work had been done on lower limb activity recognition also. Major studies of locomotion identification are summarized in Table I. Table I shows that very few studies on particular application are conducted using sEMG signal and also these studies used multiple muscles signals along with a higher number of features. Still there is lot of scope to exploit sEMG signal for the current application.

In this paper, a single muscle based continuous locomotion identification approach was explored. Here, sEMG signal from two lower limb muscles was recorded during three types of locomotion namely: Plane Walk (PW), Stair Ascending (SA), Stair Descending (SD). To acquire the sEMG signal, ADS1298 IC based sEMG interface was developed. Twelve feature vectors were constructed using 38 features (time and frequency domain) extracted from the recorded signal and classification performance of five different types of classifiers namely; Decision Tree (DT), k-Nearest Neighbor (KNN), Artificial Neural Network (ANN), Support Vector Machine (SVM) and Linear Discernment Analysis (LDA) were compared for all feature vectors. Also, a class dependent statistical based approach of feature selection was proposed and compared. All the results were compared subject-wise to analyze the variation in performance.

Rohit Gupta is with the EIED, Thapar University, Patiala, Punjab, India (e-mail: rohit.gupta@ thaapr.edu).

Ravinder Agarwal is with the EIED, Thapar University, Patiala, Punjab, India (corresponding author, phone: 01752393062, e-mail: ravinder_eeed@ thaapr.edu).

TABLE I MAJOR STUDIES ON LOCOMOTION IDENTIFICATION

Only mechanical senor						
	No. of Muscles	No. of features	Classifier	Max. acc.		
[19]	**	52	LDA	97%		
[20]	**	22	Phase Dependent LDA, QDA, LR	93%		
[21]	**	52	LDA	84.5%		
[22]	**	59	Phase dependent LDA	98.4%		
[23]	**	5	Phase dependent classifier	98%		
[24]	**	6	Locomotion dependent LDA	88%		
Only sEMG signal						
[25]	3	24	LDA, ANN, NBC	95%		
[26]	11	11	Phase dependent LDA			
[27]	12	36x45	LDA, SVM, NN	90%		
Both mechanical sensor and sEMG signal						
[28]	11	88	Phase dependent NB	100%		
[29]	7	28	LDA	97%		
[30]	9	36	Phase dependent LDA	98%		
[13]	4	60	LDA, SVM	97%		
[31]	8	86	Phase dependent SVM	95%		
[32]	9	99	Phase dependent LDA	98%		
[33]	9	124	LDA, NB	98%		
[34]	7	987	LDA, SVM	84%		

II. MATERIAL AND METHODS

A. sEMG Signal Acquisition Interface Design

The basic modules needed to design an sEMG based activity recognition system are shown in Fig. 1.

The circuit diagram to interface ADS1298 IC is shown in Fig. 2 [35]. Disposable, self-adhesive silver/silver chloride (Ag/AgCl) snap electrodes having two circular conductive areas of 1.0 cm each and an inter-electrode distance of 2.0 cm were used to pick up the electrical signal from the human body during activity. A Schottky diode protection circuit was utilized for over voltage protection and a second order RC filter of cutoff frequency 1000Hz was used for each channel. Thereafter, to amplify the signal, a programmable gain amplifier was used, with a gain of 1000. Further, the amplified signal was converted into a digital signal to transfer it to the

processing device. The amplification and analog to digital conversion was performed through one ADS1298 IC. There are many more advantages of using ADS1298 IC for such a type of system development, some of them are [36]; i) power decapitation of device goes down, ii) complexity of the system reduced, iii) cost of the system reduced, iv) due to single pin output of ADC for eight pin input, modularity of the system increased, v) sensitivity of the ADC can be varied at the programming level, vi) sampling frequency can be modified as per requirement, vii) lead off protection is provided which can be used for a safety feature, viii) programmable, adjustable gain provides the flexibility of choosing different gains.

To transfer the digital output of the ADS1298 in the processing device, an interfacing programmable device was used with the SPI protocol enabled pin. For this system, a Teensy 3.2 module was used, which is small in size and powered through the same power supply used for the ADS1298. The output (sEMG signal) of Teensy 3.2 was transferred and saved in a personal computer through serial communication that was further processed and analyzed using MATLAB 2015a.

The proposed eight channels sEMG interface cost less than 5,000 INR (approximately US\$73), whereas commercially available standard multichannel sEMG signal acquisition system costs more than 200,000 INR (approximately US\$2,900). Further, there is no need to procure any recording cables and any other interfacing device like a DAQ card or others. Here, signals were sampled at 2024 samples/sec and all other offline signal processing was done in Intel core i3 personal computer with 2.53 GHz and 3.80 GB of RAM.

B. Data Collection and Signal Processing

Five subjects having normal gait walk participated in the experimental study. Two muscles, the Fibularis longus (below the knee) M1 and Biceps Femoris (above the knee) M2 were used for the sEMG signal recording. These muscles were selected due to; ease of accessibility for the sEMG electrode placement, lower muscle mass, sustainable sEMG signal strength, etc.

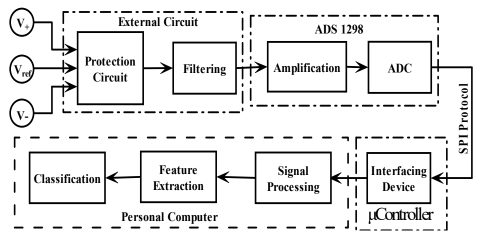


Fig. 1 Basic modules for sEMG based classification system

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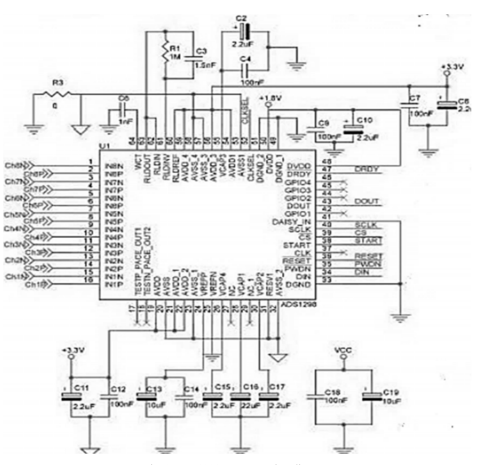


Fig. 2 ADS1298 IC connection diagram

In this study, three major locomotions of daily life were considered PW, SA and SD. Each subject was trained for five trails in each locomotion to maintain their walking speed normal and minimize the cognitive efforts. Five samples of each locomotion were recorded, each recorded sample contains five gait cycles after 20 minutes rest. To avoid muscle fatigue, 20 minutes of resting time was provided after each locomotion sample recording. The recorded signal for muscle 1 during Plain Walking (PW) is shown in Fig. 3.

The recorded sEMG signal was further processed offline; the steps needed are shown in Fig. 4. Filtering and segmentation or windowing of signal was done before extracting the features. For Filtering 4th order band pass butterworth filter, having frequency range of 60-500Hz, was applied [13].

sEMG signal was segmented in multiple windows before feature extraction (Fig. 5). Overlapped windowing techniques was applied which shows the promising results [37]–[41]. Machine learning algorithms show higher classification accuracy for window size between 200 ms to 300 ms [42], for the current research work window length of 256 ms with a shift of 32 ms was used.

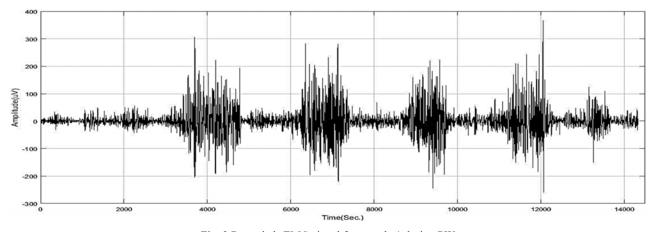


Fig. 3 Recorded sEMG signal for muscle 1 during PW

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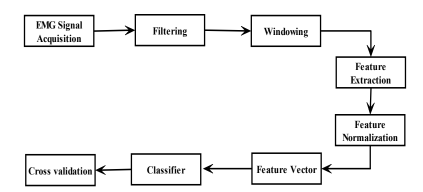


Fig. 4 Block diagram for signal processing

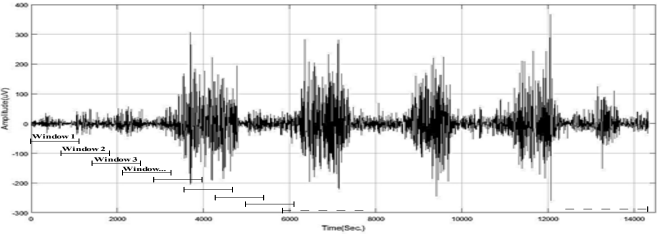


Fig. 5 Segmentation process of sEMG signal

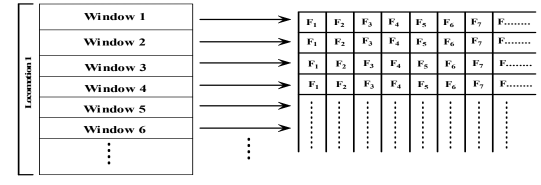


Fig. 6 Feature vector generation for each segment

Each segment of a signal was considered as a training set, the feature vector for each segment was formulated as shown in Fig. 6.

In this study, a total 38 features were extracted which includes time domain and frequency domain features presented in appendix (A1). Min-max normalization method was used to normalize the features individually to enhance the performance of classifier [43]. Further, these normalized features were grouped according to mathematical properties and information into 12 groups (Table II).

C. Classifiers and Performance Evaluation

The selection of the classifier is also an important factor

which depends on available/training data type and application. Here, five different types of classifiers, SVM, LDA, ANN, DT and KNN were compared for the application of locomotion identification.

To validate the classifier performance, repeated k-fold cross validation scheme was used. In k-fold cross validation scheme, the complete data was divided into k separate parts, out of which k-1 parts were used to train the classifier and only one part to test the classifier. To check the scheme regress, k-fold validation was repeated multiple times. Therefore, 10-fold cross validation was repeated 100 times and the average of that was used as representative performance. The performance measure was quantified as

overall % accuracy (1).

$$\% Acc = \frac{No.of \ correctly \ classified \ testing \ data}{Total \ no.of \ applied \ testing \ data} X100\%$$
(1)

TABLE II Feature Vector Information

Abv./ No of features	Features in feature vector	Vector methodology		
TD/23	All time domain features	Features having time domain information		
FD/11	All frequency domain features	Features having frequency domain information		
AR/4	AR1, AR2, AR3, AR4	Predictive information based features		
TFD/34	All time and frequency domain features	All time and frequency domain features		
TAD/27	All time and predictive domain features	Time and predictive information based features		
FAD/15	All frequency and predictive domain features	Frequency and predictive information based features		
TFAD/38	All time, frequency and predictive domain features	Time, frequency and predictive information based features		
HuF/4	MAV, WL, ZC, SSC	Hudgin's feature vector [44]		
DuF/6	IEMG, VAR, WL, ZC, SSC, WAMP	Du's feature vector [45]		
EnF/8	IEMG, MAV, MAV1, MAV2, SSI, VAR, RMS, LOG	Features having energy information		
CoF/6	WL, AAC, DASDV, TM3, 4, 5	Complexity information based features		
FITD/4	ZC, MYOP, WAMP, SSC	Features having frequency information in time domain		
CSF/16	SSI, VAR, TM3, 4, 5, DASDV, H1,2,4,5, MNF, MDF, PKF, FR, PSR, VCF, AR4	Class dependent statistical approach (proposed)		

III. PROPOSED FEATURE SELECTION ALGORITHM

The performance of any classifier is highly dependent on information used for the training of classifier. The redundant information/features were avoided as these redundant features act as noise and the classifiers suffers from over fitting. The steps involved in the proposed method (for three activities) for a class dependent statistical approach are as follows:

Step 1. Record the training data class wise.

Step 2. Extract the features from the training data.

Step 3. Consider one feature at a time:

- Arrange the data in all possible combinations/groups pair of activities (1&2, 2&3 and 3&1);
- Apply one way ANOVA analysis to check whether the feature have significant difference in between activity; and,
- iii. Store the p values of feature w.r.to activity combinations.
- Step 4. Repeat step 3 for each feature (after this step, users have three p value for each feature corresponding to pair of activities combinations).
- Step 5. Rank each feature as per their p value, the smaller the p value, the higher the ranking (after this step, users have three ranks for each feature corresponding to pair of activities combinations).
- Step 6.Select the top performing features (lower p values, p<0.001) from each group and generate feature vector

(common features in more than one group will be considered ones).

The basic methodology applied for feature selection was to rank the individual feature as per their ability to distinguish between pair of activities, for that p value was considered (*lower the p value higher the distinguishability*). To check the p value of each feature one way ANOVA analysis was performed for individual feature. Thereafter, features having p<0.001 from all pair of activities group put together to construct feature vector. Total of 16 features were selected out of 38 features and most of them are time domain features.

IV. RESULTS AND DISCUSSION

A. Effect of Subject Variability on Classification Accuracy

To determine the effect of subject variability on classification accuracy, one-way analysis of variance (ANOVA) test was performed on each classifier individually for both the muscles.

Table III shows the p-values obtained for each classifier and muscles.

TABLE III P-Values for One Way ANOVA						
Sr. No.	Classifier	Muscle 1	Muscle 2			
1.	SVM	0.85	0.50			
2. LDA		0.09	0.09			
3.	NN	0.50	0.06			
4.	DT	0.76	0.0009			
5.	KNN	0.89	0.0001			

It was observed that subject variability does not have a statistically significant effect on classifiers performance as for all the classifies the obtained p value > 0.05, only DT and KNN classifiers have significant difference in classifier performance with respect to subjects. To reduce the complexity of problem from comparison point of view, classification performance was averaged over the five subjects for further analysis.

B. Effect of Muscle Variability on Classification Accuracy

To determine the effect of muscle variability on classification accuracy, one-way analysis of variance (ANOVA) test was performed on each classifier individually. The subject variation was removed by averaging the classification performance. Over all second muscle performs better if all the feature vectors were given same weightage, as illustrated in Fig. 7.

C. Feature Vector & Classifier Performance

Table IV illustrates the classification performance of various classifiers for each feature vector. Top 50% best performing feature vectors are highlighted for each classifier. The major top performing feature vectors are TD, FD, TFD, TAD, FAD, TFAD and CSF for all the classifiers.

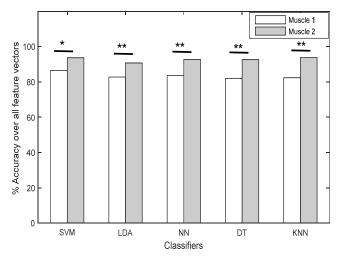


Fig. 7 Averaged classification accuracy of various classifiers for both the muscles (* *p*-value<0.01, ** *p*-value<0.001)

For the SVM classifier and muscle 1, the TD feature vector performs best (98.79 \pm 0.37), as compared to the other feature vectors, while CSF was the second best (98.73 \pm 0.68) performing feature vector. In order to test the significance of classification accuracy, ANOVA test was applied with a

significance level of 0.01. The ANOVA test shows no significant difference in performance of TD and CSF. For muscle 2, TD was the best (98.73 \pm 0.53) performing feature vector, whereas TFD, TAD were the second best (97.84 \pm 0.94), (97.84 \pm 0.72) performing feature vectors and ANOVA test showed no significant difference in performance with significance level of 0.01.

For the LDA classifier, the results show that the TAFD and TFD feature vectors performed best (99.65 \pm 0.36 for muscle 1 and 99.05 \pm 0.42 for muscle 2), and second best (98.32 \pm 0.53 for muscle 1 and 97.84 \pm 0.60 for muscle 2), and for both the muscles, the ANOVA test also showed a significant difference in classification accuracy with significance level of 0.01. For the NN classifier, the results of muscle 1 demonstrates the TFAD feature vector as the best (98.73 \pm 0.82) performing feature vector and CSF as the second best (97.08 \pm 1.97) performing feature vector with statistically significant difference with a significance level of 0.01; whereas for muscle 2, TFAD (97.94 \pm 0.63) and TD (96.89 \pm 1.02) were the best and second best performed feature vectors, but with no statistically significant difference. Similarly, the results shown for classifier DT and KNN were interpreted.

 TABLE IV

 PREDICTIVE PERFORMANCE (%CLASSIFICATION ACCURACY) OF CLASSIFIERS FOR DIFFERENT FEATURE VECTORS

	PREDICTIVE PERFORMANCE (%CLASSIFICATION ACCURACY) OF CLASSIFIERS FOR DIFFERENT FEATURE VECTORS									
	SVM		LDA		NN		DT		KNN	
	M1	M2	M1	M2	M1	M2	M1	M2	M1	M2
TD	98.79±0.37	98.73±0.53	96.29±1.06	97.14±0.49	97.05±1.35	96.89±1.02	93.78±1.26	93.87±0.52	92.06±1.35	97.02±0.52
FD	$97.05 {\pm} 0.65$	$94.98{\pm}1.14$	88.35±2.11	90.13±1.75	$82.54{\pm}7.53$	$93.14{\pm}2.88$	91.17±1.75	$93.68{\pm}1.23$	$90.41{\pm}1.14$	95.78±1.23
AR	70.16 ± 2.69	$89.87{\pm}1.73$	67.49 ± 3.52	85.71±2.79	$68.03{\pm}2.24$	$89.21{\pm}1.84$	$63.81{\pm}5.85$	$92.76{\pm}1.52$	$68.79{\pm}2.80$	92.38±1.52
TFD	96.54±1.13	$97.84{\pm}0.94$	$98.32{\pm}0.53$	$97.84{\pm}0.60$	95.62 ± 2.00	96.29±1.16	$84.13 {\pm} 5.02$	93.24±1.23	$89.21{\pm}1.82$	94.92±1.23
TAD	$96.86{\pm}0.98$	$97.84{\pm}0.72$	$96.86{\pm}0.76$	97.11 ± 0.93	95.81±1.55	$96.44{\pm}1.91$	$93.43{\pm}1.38$	$94.32{\pm}1.26$	$83.59{\pm}1.42$	92.41±1.26
FAD	$98.57 {\pm} 0.65$	95.65 ± 0.82	93.27±1.93	$90.86{\pm}1.78$	96.35 ± 2.38	94.38±1.65	$91.40{\pm}1.32$	$93.68{\pm}1.52$	$94.92{\pm}0.55$	95.05±1.52
TFAD	$97.24{\pm}1.00$	97.75±0.71	99.65 ± 0.36	99.05 ± 0.42	$98.73{\pm}0.82$	$97.94{\pm}0.63$	$90.19{\pm}1.60$	$93.78{\pm}0.86$	91.62 ± 0.92	96.03 ± 0.866
HuF	$69.84{\pm}2.00$	$88.73{\pm}1.81$	66.79 ± 3.09	86.25 ± 2.40	$68.00{\pm}2.47$	$89.02{\pm}1.80$	66.41 ± 2.34	$87.71 {\pm} 1.60$	$67.97{\pm}2.68$	91.24±1.60
DuF	74.57 ± 1.76	$89.87{\pm}1.42$	69.24 ± 3.07	$85.33{\pm}2.04$	73.37 ± 3.77	$89.56{\pm}1.78$	$71.97{\pm}1.84$	$92.95{\pm}1.21$	$75.94{\pm}1.35$	92.63±1.21
EnF	$81.90{\pm}1.46$	90.67±2.1	70.67 ± 2.26	85.46 ± 2.22	73.17 ± 5.36	89.14±2.33	91.05 ± 1.62	93.21±1.44	81.11 ± 1.70	92.57±1.44
CoF	74.57 ± 2.99	90.06 ± 2.19	$69.17 {\pm} 2.67$	$85.59{\pm}1.97$	71.46 ± 3.62	$89.59{\pm}1.67$	$72.60{\pm}2.62$	92.51±1.79	76.16 ± 2.48	92.41±1.79
FITD	$69.87{\pm}1.83$	$88.73{\pm}1.81$	67.27 ± 3.40	86.25 ± 2.4	$67.49{\pm}1.81$	$89.02{\pm}1.80$	65.65 ± 2.34	$87.71 {\pm} 1.60$	$67.84{\pm}1.74$	91.24±1.60
CSF	$98.73{\pm}0.68$	96.98±0.59	$92.38{\pm}2.20$	$92.63{\pm}1.82$	$97.08 {\pm} 1.97$	94.95±1.93	$90.70{\pm}1.96$	94.16±1.25	90.70±1.78	94.73±1.25

The results shown by the classifiers, possess quit variability in terms of best performing feature vectors. Feature vectors having a maximum number of features did not perform best for each classifier, which reflects the variability and unique nature of the classifiers. So, for each classifier, comparative analysis is necessary in order to find a suitable feature vector. One common reflection made by all classifiers was that feature vectors having time domain features performs comparatively better, which suggests one should pay more attention towards time domain based features. The feature vector (CSF) selected through the proposed feature selection algorithm stood well against other feature vectors in terms of performance and number of features. But, there is still scope for improvement in the feature selection algorithm, and the presented results suggest including the variability of the classifiers with the feature vector selection algorithm.

To identify the best performed classifier, the predictive performance of the top 50% feature vectors was averaged over the feature vector and muscles as well; the result is shown in Fig. 8. The SVM classifier possess the best accuracy as compared to other classifiers, whereas ANOVA test shows all the classifier pairs performed statistically similarly for the top 50% selected feature vectors, apart from the classifier pair SVM and DT.

To quantify the effects of experimental factors (Feature vector, Muscle and classifier), a multifactor analysis of variance (ANOVA) on classification accuracy was performed. It was observed that all three experimental factors and their interactions were the significant factors affecting the prediction performance, as their p-value was <0.01 for

significance level 0.01. Fig. 9 shows that feature vector is the most dominant source of variability with 51.51% of total variance in classification accuracy. Muscle selection also plays an important role, as it possesses 20.83% of total variance. Whereas classifier selection illustrates the least percentage variance (1.26%) among the three compared experimental factors.

V.CONCLUSION

In this paper, a low cost multichannel sEMG interface for clinical and research laboratory purpose was developed. The developed interface utilized ADS1298 integrated circuit, which reduces the problem of existing interfaces like high cost, power combustion and complex circuitry burden. It provides the flexibility to modify the gain of amplifier and sensitivity of ADC without making any change in the circuitry. Further, to prove the suitability and applicability of the developed interface, it was used to acquire the lower limb sEMG signal from two muscles for three locomotions. A comparative study is presented for locomotion identification. The study provides a regress comparison of multiple factors like feature vectors, muscle, classifiers along with class dependent statistical approach of feature selection. The presented results revel that muscle 2 Biceps Femoris (above the knee) performs better for locomotion identification. Also, time domain based features prove their suitability for current application, while multifactor analysis of variance (ANOVA) analysis suggests that classifier selection has least variability in terms of classification accuracy, whereas feature vector selection possesses the most. The potential outcomes of the presented research work will be useful in future for prosthesis control.

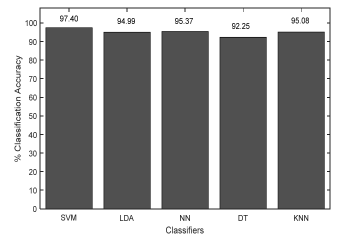


Fig. 8 Classification accuracy for different classifiers over selected feature vectors

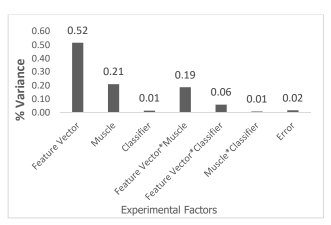


Fig. 9 Variance of classification accuracy shared by experimental factors and their interaction

	TABLE V						
		NAME AND DEFINITION OF EXT	RACTED FEATURES [9]				
Sr. No.	Abv.	Name of Feature	Definition				
1	IEMG	Integrated EMG	$IEMG = \sum_{i=1}^{N} X_i $				
2	MAV	Mean absolute value	$MAV = \frac{1}{N} \sum_{i=1}^{L-N} X_i $				
3	MAV1	Modified mean absolute value type 1	$MAV1 = \frac{1}{N} \sum_{i=1}^{N} w_i X_i , w_i = \begin{cases} 1 & \text{if } 0.25N \le i \le 0.75N \\ 0.5 & \text{otherwise} \\ (1 & \text{if } 0.25N \le i \le 0.75N) \end{cases}$				
4	MAV2	Modified mean absolute value type 2	$MAV1 = \frac{1}{N} \sum_{i=1}^{N} w_i X_i , w_i = \begin{cases} i=1 \\ 1 \text{ if } 0.25N \le i \le 0.75N \\ 0.5 \text{ otherwise} \end{cases}$ $MAV2 = \frac{1}{N} \sum_{i=1}^{N} w_i X_i , w_i = \begin{cases} 1 \text{ if } 0.25N \le i \le 0.75N \\ \frac{4i}{N} \text{ elseif } i < 0.25N \\ \frac{4(i-N)}{N} \text{ otherwise} \end{cases}$				
5	SSI	Simple square integral	$SSI = \sum_{i=1}^{N} X_i^2$				
6	VAR	Variance of EMG	$VAR = \frac{1}{N-1} \sum_{i=1}^{N} X_i^2$				
7	TM3	Absolute value of the 3 rd temporal moment	$TM3 = \left \frac{1}{N} \sum_{i=1}^{N} X_i^3 \right $				
8	TM4	Absolute value of the 4 th temporal moment	$TM4 = \left \frac{1}{N} \sum_{i=1}^{N} X_i^4 \right $				

APPENDIX TABLE V Name and Definition of Extracted Features [9]

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9TMSAbsolute value of the 5th temporal moment $TMS = \left \frac{1}{2k} \sum_{i=1}^{N} K_i^i \right $ 10RMSRoot mean square $RMS = \int_{1}^{N} \sum_{i=1}^{N} K_i^i$ 11LOGLog detector $LOG = e^{\frac{1}{2} \sum_{i=1}^{N} M_i^i}$ 12WLWaveform length $WL = \sum_{i=1}^{N-1} K_{i+1} - K_i $ 13AACAverage amplitude change $AAC = \frac{1}{N} \sum_{i=1}^{N-1} K_{i+1} - K_i $ 14DASDVDifference absolute standard deviation value $DASDV = \int_{1}^{1} \sum_{i=1}^{N-1} (K_{i+1} - K_i)^2$ 15ZCZero crossing $ZC = \sum_{i=1}^{N-1} [sgn(X_iXX_{i+1}) \cap [X_i - X_{i+1}] \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N-1} (K_{i+1} - K_{i+1}) \ge threshold]$ 17WAMPWillison amplitude $WAMP = \sum_{i=1}^{N-1} [f(X_i - X_{i-1})X_i - X_{i+1}] \ge threshold]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i-1})X_i(X_i - X_{i+1})]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i-1})X_i - X_{i+1})]$ 19H1H1H121H3H1 stogram of EMG23H5H424MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11" Spectral moments30SM22" Spectral moments31SM33" Spectral moments32FRFrequency ratio33PSRPo	Sr. No.	Abv.	Name of Feature	Definition
10RMSRoot mean square $RMS = \left \frac{1}{N} \sum_{l=1}^{N} X_{l}^{2} \right $ 11LOGLog detector $LOG = e^{\frac{1}{N} \sum_{l=1}^{N} X_{l+1} - X_{l} }$ 12WLWaveform length $WL = \sum_{l=1}^{l=1} X_{l+1} - X_{l} $ 13AACAverage amplitude change $AAC = \frac{1}{N} \sum_{l=1}^{l=1} X_{l+1} - X_{l} $ 14DASDVDifference absolute standard deviation value $DASDV = \int \prod_{l=1}^{l-1} X_{l+1} - X_{l} $ 15ZCZero crossing $ZC = \sum_{l=1}^{l=1} [Sgn(X_l \times X_{l+1}) \cap X_l - X_{l+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MTOP = \frac{1}{N} \sum_{l=1}^{l} I_{l}(X_{l}) - X_{l+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MTOP = \frac{1}{N} \sum_{l=1}^{l} I_{l}(X_{l}) - X_{l+1} \ge threshold]$ 17WAMPWillison amplitude $MTOP = \sum_{l=1}^{l} I_{l}(X_{l}) - X_{l+1} \ge threshold]$ 18SSCSlope sign change $SSC = \sum_{l=1}^{l} [I_{l}(X_{l} - X_{l-1}) \times (X_{l} - X_{l+1})]$ 18SSCSlope sign change $SSC = \sum_{l=1}^{l} [I_{l}(X_{l} - X_{l-1}) \times (X_{l} - X_{l+1})]$ 19H110 $IK2$ 21H3Histogram of EMGHistogram of EMG23H524MNFMean frequency24MNFMean frequency $NF = \sum_{l=1}^{l} p_{l} - \sum_{l=1}^{l} p_{l}$				
11LOGLog detectorLOG = $e^{\frac{1}{N} \sum_{i=1}^{N} \log(N_i)}$ 12WLWaveform length $WL = \sum_{i=1}^{N} X_{i+1} - X_i $ 13AACAverage amplitude change $AAC = \frac{1}{N} \sum_{i=1}^{N-1} X_{i+1} - X_i $ 14DASDVDifference absolute standard deviation value $DASDV = \sqrt{\prod_{i=1}^{N-1} \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2}$ 15ZCZero crossing $ZC = \sum_{i=1}^{N-1} [sgn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N} [f(X_i)]$ 17WAMPWillison amplitude $MXP = \sum_{i=1}^{N-1} [f(X_i - X_{i+1})] \ge threshold]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 19H1 $G(X) = \left\{ 1, if X \ge threshold \\ 0, otherwise \end{bmatrix}$ 24MNFMean frequency $MNF = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 25MDFMedian frequency $MNP = \sum_{i=1}^{N-1} p_i - \sum_{i=1}^{N} p_i$ 26PKFPeak frequency $MNP = \sum_{i=1}^{N-1} p_i - \sum_{i=1}^{N} p_i$ 25MDFMean power $MNP = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 26PKFPeak frequency $MNP = \sum_{i=1}^{N-1} p_i$ 27MNPMean power $MNP = \sum_{i=1}^{N} p_i - SMO$ 29SM11" Spectral moments $SM = \sum_{i=1}^{N} p_i - SMO$ 30SM2 2^{ni} Spectral moments $SM = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 31SM33" Spectral mome	9	TM5	Absolute value of the 5 th temporal moment	$TM5 = \left \frac{1}{N} \sum_{i=1}^{X_i^5} \right $
11LOGLog detectorLOG = $e^{\frac{1}{N} \sum_{i=1}^{N} \log(N_i)}$ 12WLWaveform length $WL = \sum_{i=1}^{N} X_{i+1} - X_i $ 13AACAverage amplitude change $AAC = \frac{1}{N} \sum_{i=1}^{N-1} X_{i+1} - X_i $ 14DASDVDifference absolute standard deviation value $DASDV = \sqrt{\prod_{i=1}^{N-1} \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2}$ 15ZCZero crossing $ZC = \sum_{i=1}^{N-1} [sgn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N} [f(X_i)]$ 17WAMPWillison amplitude $MXP = \sum_{i=1}^{N-1} [f(X_i - X_{i+1})] \ge threshold]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 18SSCSlope sign change $SSC = \sum_{i=1}^{N-1} [f(X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 19H1 $G(X) = \left\{ 1, if X \ge threshold \\ 0, otherwise \end{bmatrix}$ 24MNFMean frequency $MNF = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 25MDFMedian frequency $MNP = \sum_{i=1}^{N-1} p_i - \sum_{i=1}^{N} p_i$ 26PKFPeak frequency $MNP = \sum_{i=1}^{N-1} p_i - \sum_{i=1}^{N} p_i$ 25MDFMean power $MNP = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 26PKFPeak frequency $MNP = \sum_{i=1}^{N-1} p_i$ 27MNPMean power $MNP = \sum_{i=1}^{N} p_i - SMO$ 29SM11" Spectral moments $SM = \sum_{i=1}^{N} p_i - SMO$ 30SM2 2^{ni} Spectral moments $SM = \sum_{i=1}^{N} p_i - \sum_{i=1}^{N} p_i$ 31SM33" Spectral mome	10	RMS	Root mean square	$RMS = \int \frac{1}{N} \sum_{i=1}^{N} X_i^2$
13AACAverage amplitude change $AAC = \frac{1}{N} \sum_{i=1}^{N-1} X_{i+1} - X_i $ 14DASDVDifference absolute standard deviation value $DASDV = \sqrt{1 - \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2}$ 15 ZC Zero crossing $ZC = \sum_{i=1}^{N-1} Syn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N} [f(X_i))$ 17WAMPWillison amplitude $MAP = \sum_{i=1}^{N-1} [f(X_i - X_{i+1})]$ 18SSCSlope sign change $SSC = \sum_{i=2}^{N-1} [f(X_i - X_{i+1})]]$ 18SSCSlope sign change $f(X) = \left\{ 1.if X \ge threshold \\ 0. otherwise$ 19H1Distormation $SSC = \sum_{i=2}^{N-1} [f(X_i - X_{i+1})]]$ 18SSCSlope sign change $f(X) = \left\{ 1.if X \ge threshold \\ 0. otherwise$ 19H1Distormation $SSC = \sum_{i=2}^{N-1} [f^P_i] / \sum_{i=1}^{N} P_i]$ 21H3Histogram of EMGHistogram of EMG23H5 MNF Mean frequency24MNFMean frequency $MNF = \sum_{i=1}^{M} P_i = \frac{1}{2} \sum_{i=1}^{N} P_i$ 25MDFMedian frequency $MNF = \sum_{i=1}^{M} P_i = \frac{1}{2} \sum_{i=1}^{M} P_i$ 26PKFPeak frequency $PKF = max(P_i), j = 1, 2,, M$ 27MNPMean power $MNP = \sum_{i=1}^{N} P_i$ 28TTPTotal power $TTP = \sum_{i=1}^{N} P_i$ 30SM2 2^{nd} Spectral moments $SM2 = \sum_{i=1}^{N} P_i f_i^2$ 31SM3 3^{nd} Spectral moments $SM3 = \sum_{i=1}^{N} P_i f_i^2$ 32FR <td>11</td> <td>LOG</td> <td>Log detector</td> <td>N</td>	11	LOG	Log detector	N
14DASDVDifference absolute standard deviation value $DASDV = \sqrt{\frac{1}{N-1}} \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2$ 15ZCZero crossing $ZC = \sum_{i=1}^{N-1} [sgn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N} [f_i(X_i)]$ 16MYOPWyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N-1} [f_i(X_i)]$ 17WAMPWillison amplitude $MXMP = \sum_{i=1}^{N-1} [f_i(X_i - X_{n+1})]$ 18SSCSlope sign change $SSC = \sum_{i=2}^{N-1} [f_i(X_i - X_{n+1})]$ 19H1 $f(X) = \left(1, if X \ge threshold \\ 0, otherwise$ 20H2 $H1$ 21H3Histogram of EMG23H5Histogram of EMG24MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11" Spectral moments30SM22 nd Spectral moments31SM33 rd Spectral moments32FRFrequency ratio33PSRPower spectrum ratio	12	WL	Waveform length	$WL = \sum_{i=1}^{N} X_{i+1} - X_i $
14DASDVDifference absolute standard deviation value $DASDV = \sqrt{\frac{1}{N-1}} \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2$ 15ZCZero crossing $ZC = \sum_{i=1}^{N-1} [sgn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$ 16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N} [f_i(X_i)]$ 16MYOPWyopulse percentage rate $MYOP = \frac{1}{N} \sum_{i=1}^{N-1} [f_i(X_i) - X_{i+1}] \ge threshold$ 17WAMPWillison amplitude $WAMP = \sum_{i=1}^{N-1} [f_i(X_i - X_{i+1})]$ 18SSCSlope sign change $SSC = \sum_{i=2}^{N-1} [f_i(X_i - X_{i+1})]$ 19H1 $f(X) = \left\{ 1.if_i X \ge threshold \\ 0.otherwise$ 20H2 $H1$ $H1$ 21H3Histogram of EMG23H5H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11" Spectral moments30SM22 nd Spectral moments31SM33 nd Spectral moments32FRFrequency ratio33PSRPower spectrum ratio	13	AAC	Average amplitude change	$AAC = \frac{1}{N} \sum_{i=1}^{N-1} X_{i+1} - X_i $
16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{j=1}^{N} [f(X_i)]$ 17WAMPWillison amplitude $f(X) = \begin{cases} 1, if X \ge threshold \\ 0, otherwise \end{cases}$ 17WAMPWillison amplitude $WAMP = \sum_{i=1}^{M} [f(X_i - X_{i+1})]$ 18SSCSlope sign change $SSC = \sum_{i=2}^{N-1} [f((X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 18SSCSlope sign change $f(X) = \begin{cases} 1, if X \ge threshold \\ 0, otherwise \end{cases}$ 19H1 $SSC = \sum_{i=2}^{N-1} [f(X_i - X_{i-1}) \times (X_i - X_{i+1})]]$ 20H2 $f(X) = \{1, if X \ge threshold \\ 0, otherwise \end{cases}$ 21H3Histogram of EMG22H4 $\sum_{i=1}^{M} P_i - \sum_{i=1}^{M} P_i = \sum_{i=1}^{M} P_i$ 24MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNP28TTP29SM11 st Spectral moments30SM22 ^{sts} Spectral moments31SM33 st Spectral moments32FRFrequency ratio33PSRPower spectrum ratio9PSR $P_{i} = \sum_{i=1}^{M} P_{i}$	14	DASDV	Difference absolute standard deviation value	$DASDV = \sqrt{\frac{1}{N-1} \sum_{i=1}^{N-1} (X_{i+1} - X_i)^2}$
16MYOPMyopulse percentage rate $MYOP = \frac{1}{N} \sum_{j=1}^{N} [f(X_i)]$ 17WAMPWillison amplitude $f(X) = \begin{cases} 1, if X \ge threshold \\ 0, otherwise \end{cases}$ 17WAMPWillison amplitude $WAMP = \sum_{i=1}^{M} [f(X_i - X_{i+1})]$ 18SSCSlope sign change $SSC = \sum_{i=2}^{N-1} [f((X_i - X_{i+1}) \times (X_i - X_{i+1})]]$ 18SSCSlope sign change $f(X) = \begin{cases} 1, if X \ge threshold \\ 0, otherwise \end{cases}$ 19H1 $SSC = \sum_{i=2}^{N-1} [f(X_i - X_{i-1}) \times (X_i - X_{i+1})]]$ 20H2 $f(X) = \{1, if X \ge threshold \\ 0, otherwise \end{cases}$ 21H3Histogram of EMG22H4 $\sum_{i=1}^{M} P_i - \sum_{i=1}^{M} P_i = \sum_{i=1}^{M} P_i$ 24MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNP28TTP29SM11 st Spectral moments30SM22 ^{sts} Spectral moments31SM33 st Spectral moments32FRFrequency ratio33PSRPower spectrum ratio9PSR $P_{i} = \sum_{i=1}^{M} P_{i}$	15	ZC	Zero crossing	$ZC = \sum_{i=1}^{N-1} [sgn(X_i \times X_{i+1}) \cap X_i - X_{i+1} \ge threshold]$
$f(X) = \begin{cases} I, If X \ge threshold \\ 0, otherwise \\ WAMP = \sum_{i=1}^{N-1} [f(X_n - X_{n+1})] \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ WAMP = \sum_{i=1}^{N-1} [f(X_n - X_{n+1})] \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ SSC = \sum_{i=2}^{N-1} [I[(X_i - X_{i-1}) \times (X_i - X_{i+1})]] \\ 18 SSC Slope sign change \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ I = I \\ I = $				$sgn(X) = \begin{cases} 1, if \ X \ge thrsehold \\ 0, otherwise \end{cases}$
$f(X) = \begin{cases} I, If X \ge threshold \\ 0, otherwise \\ WAMP = \sum_{i=1}^{N-1} [f(X_n - X_{n+1})] \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ WAMP = \sum_{i=1}^{N-1} [f(X_n - X_{n+1})] \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ SSC = \sum_{i=2}^{N-1} [I[(X_i - X_{i-1}) \times (X_i - X_{i+1})]] \\ 18 SSC Slope sign change \\ f(X) = \{ I, If X \ge threshold \\ 0, otherwise \\ I = I \\ I = $	16	МҮОР	Myopulse percentage rate	$MYOP = \frac{1}{N} \sum_{i=1}^{N} [f(X_i)]$
18SSCSlope sign change $SSC = \sum_{l=2}^{N-1} [f[(X_l - X_{l-1}) \times (X_l - X_{l+1})]]$ 18SSCSlope sign change $f(X) = \left\{ \substack{1, if \ X \ge threshold \\ 0, otherwise} \right\}$ 19H120H221H3Histogram of EMG22H423H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11 st Spectral moments30SM22 nd Spectral moments31SM33 rd Spectral moments32FRFrequency ratio33PSRPower spectrum ratio				$f(X) = \begin{cases} 1, & \text{if } X \ge \text{threshold} \\ 0, & \text{otherwise} \end{cases}$
18SSCSlope sign change $SSC = \sum_{l=2}^{N-1} [f[(X_l - X_{l-1}) \times (X_l - X_{l+1})]]$ 18SSCSlope sign change $f(X) = \left\{ \substack{1, if \ X \ge threshold \\ 0, otherwise} \right\}$ 19H120H221H3Histogram of EMG22H423H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11 st Spectral moments30SM22 nd Spectral moments31SM33 rd Spectral moments32FRFrequency ratio33PSRPower spectrum ratio	17	WAMP	Willison amplitude	$WAMP = \sum_{\substack{i=1\\(1,i\neq K\}}} [f(X_n - X_{n+1})]$
18SSCSlope sign changeInterference19H120H221H3Histogram of EMG22H423H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11" Spectral moments30SM22" Spectral moments31SM33" Spectral moments32FRFrequency ratio33PSRPower spectrum ratio				$f(X) = \begin{cases} 1, l & X \ge lineshold \\ 0, otherwise \end{cases}$
19H120H221H3Histogram of EMG22H423H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM1I* Spectral moments30SM2 2^{nd} Spectral moments31SM3 3^{nd} Spectral moments32FRFrequency ratio33PSRPower spectrum ratioPSR $\frac{P_0}{P} = \frac{\sum_{j=1}^{U_{j=1}} P_j}{\sum_{j=1}^{J_{j=1}} P_j}$	18	SSC	Slope sign change	$SSC = \sum_{i=2}^{\infty} [f[(X_i - X_{i-1}) \times (X_i - X_{i+1})]]$
20H221H3Histogram of EMG22H423H524MNFMean frequency25MDFMedian frequency26PKFPeak frequency27MNPMean power28TTPTotal power29SM11st Spectral moments30SM22nd Spectral moments31SM33rd Spectral moments32FRFrequency ratio33PSRPower spectrum ratio				$f(X) = \begin{cases} 1, & \text{if } X \ge threshold \\ 0, & \text{otherwise} \end{cases}$
21H3Histogram of EMGHistogram of EMG22H4123H524MNFMean frequency $MNF = \sum_{j=1}^{M-1} f_j P_j / \sum_{j=1}^{M} P_j$ 25MDFMedian frequency $\sum_{j=1}^{MDF} P_j = \sum_{j=MDF}^{M} P_j = \frac{1}{2} \sum_{j=1}^{M} P_j$ 26PKFPeak frequency $PKF = \max(P_j), j = 1, 2,, M$ 27MNPMean power $MNP = \sum_{j=1}^{M-1} P_j$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j$ 30SM2 2^{nd} Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM3 3^{rd} Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \sum_{j=1}^{M} P_j f_j^2$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{N-1} P_j}{\sum_{j=0}^{N-1} P_j}$	19	H1		
22H423H524MNFMean frequency $MNF = \sum_{j=1}^{M} f_j P_j / \sum_{j=1}^{H} P_j$ 25MDFMedian frequency $\sum_{j=1}^{MDF} P_j = \frac{1}{2} \sum_{j=1}^{M} P_j$ 26PKFPeak frequency $PKF = \max(P_j), j = 1, 2,, M$ 27MNPMean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SMO$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j^2$ 30SM22nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33nd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^2$ 32FRFrequency ratio $FR = \sum_{j=1L}^{M} P_j f_j^2$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{J_{j=0}} P_j}{\sum_{j=0}^{J_{j=0}} P_j}$		H2		
23H524MNFMean frequency $MNF = \sum_{j=1}^{M} f_j P_j / \sum_{j=1}^{M} P_j$ 25MDFMedian frequency $\sum_{j=1}^{MDF} P_j = \sum_{j=1}^{M} P_j = \frac{1}{2} \sum_{j=1}^{M} P_j$ 26PKFPeak frequency $PKF = \max(P_j), j = 1, 2,, M$ 27MNPMean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j^2$ 30SM22nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \sum_{j=1}^{MLL} P_j / \sum_{j=1}^{MH} P_j$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{j=1} R_j}{\sum_{j=0}^{N} R_j}$			Histogram of EMG	Histogram of EMG
24MNFMean frequency $MNF = \sum_{j=1}^{M} f_j P_j / \sum_{j=1}^{M} P_j$ 25MDFMedian frequency $\sum_{j=1}^{MDF} P_j = \sum_{j=MDF}^{M} P_j = \frac{1}{2} \sum_{j=1}^{M} P_j$ 26PKFPeak frequency $PKF = \max(P_j), j = 1, 2, \dots, M$ 27MNPMean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j$ 30SM22st Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33st Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^2$ 32FRFrequency ratio $FR = \sum_{j=1L}^{M} P_j / \sum_{j=1L}^{M} P_j$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \sum_{j=0}^{\sum_{j=0}^{M} n} P_j$				
25 MDF Median frequency $\sum_{j=1}^{MDF} P_j = \sum_{j=MDF}^{M} P_j = \frac{1}{2} \sum_{j=1}^{M} P_j$ 26 PKF Peak frequency $PKF = \max(P_j), j = 1, 2,, M$ 27 MNP Mean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28 TTP Total power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29 SM1 1 st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j^2$ 30 SM2 2 nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31 SM3 3 rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32 FR Frequency ratio $FR = \frac{\sum_{j=1}^{M} P_j f_j^3}{P_j f_j^2 = \sum_{j=1}^{M} P_j}$	23	H5		
26PKFPeak frequency $j=1$ $j=MDF$ $j=1$ 27MNPMean power $PKF = \max(P_j), j = 1, 2,, M$ 27MNPMean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j$ 30SM22nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \sum_{j=1L}^{VILL} P_j / \sum_{j=LL}^{VIH} P_j$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{f_0 - n} P_j}{\sum_{j=0}^{\infty} P_j}$	24	MNF	Mean frequency	MDF M M
27MNPMean power $MNP = \frac{\sum_{j=1}^{M} P_j}{M}$ 28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j$ 30SM22nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \frac{\sum_{j=1}^{UL} P_j}{N} / \sum_{j=1}^{ULH} P_j}$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{f_0+n} P_j}{\sum_{j=-\infty}^{S} P_j}$	25	MDF		J=1 $J=MDF$ $J=1$
28TTPTotal power $TTP = \sum_{j=1}^{M} P_j = SM0$ 29SM11st Spectral moments $SM1 = \sum_{j=1}^{M} P_j f_j$ 30SM22nd Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM33rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \frac{\sum_{j=1LL}^{M} P_j}{\sum_{j=1LL}^{N} P_j}$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=0}^{f_0+n} P_j}{\sum_{j=-\infty}^{f_0+n} P_j}$	26	PKF	Peak frequency	
29 SM1 1 st Spectral moments 30 SM2 2 nd Spectral moments 31 SM3 3 rd Spectral moments 32 FR Frequency ratio 33 PSR Power spectrum ratio J^{J-1} $SM1 = \sum_{j=1}^{M} P_j f_j^2$ $SM2 = \sum_{j=1}^{M} P_j f_j^3$ $FR = \sum_{j=1}^{M} P_j f_j^3$	27	MNP	Mean power	
30SM2 2^{nd} Spectral moments $SM2 = \sum_{j=1}^{M} P_j f_j^2$ 31SM3 3^{rd} Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32FRFrequency ratio $FR = \frac{\sum_{j=LL}^{UL} P_j}{\sum_{j=LL}^{N} P_j} \frac{1}{\sum_{j=1}^{G_n + n} P_j}$ 33PSRPower spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=1}^{G_n + n} P_j}{\sum_{j=-\infty}^{G_n - n} P_j}$	28	TTP	Total power	$TTP = \sum_{j=1}^{N} P_j = SM0$
31 SM3 3 rd Spectral moments $SM3 = \sum_{j=1}^{M} P_j f_j^3$ 32 FR Frequency ratio $FR = \frac{\sum_{j=1}^{UL} P_j}{\sum_{j=LL}^{P_L} P_j}$ 33 PSR Power spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=f_0-n}^{f_0+n} P_j}{\sum_{j=-\infty}^{\infty} P_j}$	29	SM1	1 st Spectral moments	
33 PSR Power spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=f_0-n} F_j}{\sum_{j=-\infty}^{\infty} P_j}$	30	SM2	2 nd Spectral moments	$SM2 = \sum_{j=1}^{M} P_j f_j^2$
33 PSR Power spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=f_0-n} F_j}{\sum_{j=-\infty}^{\infty} P_j}$	31	SM3	3 rd Spectral moments	$SM3 = \sum_{i=1}^{M} P_j f_j^3$
33 PSR Power spectrum ratio $PSR = \frac{P_0}{P} = \frac{\sum_{j=f_0-n} F_j}{\sum_{j=-\infty}^{\infty} P_j}$	32	FR	Frequency ratio	$FR = \frac{\sum_{j=LL}^{J-1} P_j}{\sum_{j=LH}^{J-1} P_j}$
34 VCF Variance of central frequency $VCF = \frac{1}{SM0} \sum_{j=1}^{M} P_j (f_j - f_c)^2 = \frac{SM2}{SM0} - \left(\frac{SM1}{SM0}\right)^2$	33	PSR	Power spectrum ratio	$PSR = \frac{P_0}{P} = \frac{\sum_{j=f_0-n} P_j}{\sum_{j=-\infty}^{\infty} P_j}$
	34	VCF	Variance of central frequency	$VCF = \frac{1}{SM0} \sum_{j=1}^{M} P_j (f_j - f_c)^2 = \frac{SM2}{SM0} - \left(\frac{SM1}{SM0}\right)^2$

Sr. No.	Abv.	Name of Feature	Definition
35	AR1		p
36	AR2	Auto-regressive coefficients	$X_i = \sum a_p X_{i-p} + w_i$
37	AR3	Auto-regressive coefficients	p=1
38	AR4		p is order of the AR model, w_i is white noise error term

where X_i represents the EMG signal in a segment *i* and *N* denotes length of EMG signal, f_j is frequency of the spectrum at frequency bin *j*, P_j is the EMG power spectrum at frequency bin *j* and *M* is length of the frequency bin. UL, LL: upper and lower cut-off frequency of lower frequency band UH, LH: upper and lower cut-off frequency of upper frequency band.

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