Auditory brainstem functioning in individuals with misophonia

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AUDITORY BRAINSTEM FUNCTIONING IN INDIVIDUALS WITH MISOPHONIA

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Prashanth Prabhu was involved in concept development and study design, stimulus preparation, and writing the manuscript.

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Data availability statement

The datasets used for the current study are available from the corresponding author upon reasonable request.

Conflicts of Interest: There is no conflict of interest to disclose.

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1	AUDITORY BRAINSTEM FUNCTIONING IN INDIVIDUALS WITH MISOPHONIA
2	
3	ABSTRACT
4	Purpose
5	Misophonia is not investigated much from an audiological perspective. Our study aims to
6	examine the processing of the auditory retro-cochlear pathways in individuals with misophonia.
7	Methods
8	A cross-sectional study was conducted among university students who had misophonia. The
9	revised Amsterdam Misophonia Scale was used to determine the severity of misophonia.
LO	Participants were divided into mild and moderate-severe misophonia and compared with the
l1	healthy control group. Auditory Brainstem Response testing was recorded from all the
12	individuals with misophonia. The absolute latency, amplitude, inter-peak latency difference, and
13	inter-rate latency difference were compared between the groups.
L4	Results
L 5	One-way ANOVA result showed no significant difference in all the parameters of auditory
L6	brainstem response between the groups. These results are suggestive of normal brainstem
L7	processing in individuals with misophonia.
18	Conclusions
19	The study concludes that the auditory pathway up to brainstem areas is intact in individuals with
20	misophonia. Further studies are essential on a larger population for generalizing the results.
21	Keywords : Misophonia; brainstem pathway; Brainstem response; Neurophysiology; Audiology
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23	
24	

1.0 INTRODUCTION

27	Misophonia is a disorder characterized by decreased sound tolerance to specific sound stimuli,
28	known as triggers (Swedo et al., 2022). The triggers can be visual, auditory, or motor, and they
29	may cause both emotional and physiological reactions, including anxiety, increased heart rate,
30	sweating, rage, and irritation. The prevalence of misophonia is high, ranging from 23.28% (Aryal
31	& Prabhu., 2022) to 49.1% (Naylor et al., 2021). Geographical region, variation in the sample,
32	and methodology used could be the factors for variation in the prevalence rate.
33	Misophonia has not been categorized as a separate disorder by the Diagnostic and statistical
34	manual (DSM-V) of mental disorders. This disorder borders neurology, physiology, and
35	audiology (Danesh & Aazh., 2020), which can alter the physiological mechanism of the sufferers
36	and result in distraction and annoyance. The lack of separate categorization hinders the
37	recognition of the team members involved in the assessment and management of misophonia and
38	hinders the sufferers from seeking help.
39	From the audiological perspective, less research is done to understand misophonia. Various
40	researchers have shown abnormal activation of the cortical auditory structures in individuals with
41	misophonia (Kumar et al., 2021; Kumar et al., 2017; Grossini et al., 2022) through radiological
42	investigations. In addition, studies have shown abnormal processing of the autonomic nervous
43	system, including the limbic system, among misophonic (Kumar et al., 2017). However, no
44	studies have reported abnormal neural processing of the retro-cochlear structures, including
45	auditory nerve and brainstem structures, through radiological and audiological investigations.
46	The pathophysiology of misophonia is not clear yet. However, various mechanism, origin,
47	theories, and model has been explained in the literature to explain misophonia (Grossini et al.,
48	2022; Aryal & Prabhu., 2022). To understand misophonia from an audiological perspective, a
49	new model has been developed which shows the linkage of misophonia with the classical and
50	non-classical auditory pathway (Aryal & Prabhu., 2022).
51	Auditory evoked potentials (AEP) examine the auditory nerve fibers' synchronous discharge and
52	detect abnormal neuronal activation. The waveform within the few ten milliseconds of the
53	auditory evoked potentials is called auditory brainstem response (ABR). Auditory brainstem
54	response (ABR) is the electrophysiological test that measures the neural activation of the
55	auditory pathway from the dorsal auditory nerve to the inferior colliculus (Picton et al., 2000).

56	Out of seven peaks of ABR, the first, third, and fifth peaks are clinically significant and arise
57	from the dorsal part of the auditory nerve, cochlear nucleus, and lateral lemniscus, respectively
58	(Picton & Durieux Smith., 1978). The shift in the latency of the Vth peak by greater than 0.8
59	seconds, when the rate increased from 11.1/sec to 90.1/sec, indicates retro cochlear pathology
60	(Picton and Durieux Smith., 1978).
61	
62	The primary aim of our study is to evaluate the processing of the retro-cochlear pathway in
63	individuals with misophonia through electrophysiological tests. Abnormal ABR is reported
64	among individuals with sound disorders such as tinnitus and hyperacusis (Sand and Saunte,
65	1994). Misophonia is a disorder that occurs in isolation or with other sound disorders such as
66	tinnitus, hyperacusis, and phonophobia. We can hypothesize that abnormal neural processing up
67	to the brainstem level might occur among individuals with misophonia due to its co-existence
68	with these disorders. Hence, we aim to evaluate the peripheral retro-cochlear auditory pathway in
69	individuals with misophonia by administering ABR testing. The present study was designed to
70	determine whether significant differences exist in the ABR parameters in individuals with
71	misophonia.
72	
73	2.0 METHODS
74	The All India Institute of Speech and Hearing institutional ethical review board reviewed the
75	study protocol, and the ethical approval number was SH/ERB/2022-24/37. All the participants
76	were informed about the study procedure before enrolling, and written informed consent was
77	taken from all the participants.
78	2.1 Study participants
79	The cross-sectional study was conducted among individuals with clinically significant
80	misophonia, and a comparison was made with the healthy control group. To find the prevalence
81	and severity of misophonia, the survey was conducted among the students of Mysore University
82	using the Revised Amsterdam misophonia questionnaire (Jager et al., 2020). The survey invited
83	30 individuals with misophonia symptoms to the study. All the participants were 18 to 40 years

84	old, with a mean age of 25 years (SD= 7.8). Most participants, 36 (90%), were female, and 4
85	(10%) were male in the misophonia group. All participants had normal hearing sensitivity in the
86	conventional pure-tone and high-frequency audiometry ranges. The control group consisted of 15
87	participants aged 20 to 40 years (Mean age= 24 years, SD= 6 years). The age and gender ratio
88	was matched to the misophonia group. All the control group participants have normal
89	audiograms in both conventional and high-frequency ranges. All the participants in the
90	misophonia group had a history of misophonia for at least three years without any psychiatric
91	and auditory disorders co-morbidities. In addition, only participants without any middle ear,
92	cardiovascular, or neurologic illness, no history of acoustic trauma, and no ototoxic medication
93	were included in both groups. Participants with hearing loss and other psychiatric and neurologic
94	co-morbidities were excluded from the study.
95	
96	2.2 Apparatus and Procedure
97	A detailed physical examination was done, including case history, otoscopic examination, and
98	general health examination. The revised Amsterdam misophonia questionnaire has been used to
99	categorize misophonia into different severity ranges (Jager et al., 2020). The questionnaire has
100	ten questions with a score ranging from 0 to 40. The score 0-10 are considered subclinical
101	misophonia symptoms, the score of 11-20 are rated as mild misophonia, 21-30 as moderate to
102	severe misophonia, and 31-40 as severe to extreme (Jager et al., 2020).
103	A hearing assessment was done using a Garson Stadler audio star pro using ANSI guidelines
104	(Frank., 1997). The supraaural-49 headphone was used for the air conduction testing of the
105	conventional pure tone audiometry, and the Radio-ear B-71 bone vibrator was used for the bone
106	conduction testing. Similarly, Sennheiser circumaural HDA200 headphone was used for high-
107	frequency audiometry. All the audiological tests were done in the soundproof room following the
108	ANSI guidelines (Frank., 1997).
109	The frequencies from 250 Hz to 8 kHz were taken to determine the Air conduction threshold.
110	Similarly, the frequencies from 250Hz to 4 kHz were taken for the bone conduction testing. The
111	four frequency averages of 500Hz, 1 kHz, 2 kHz, and 4 kHz were taken to determine the

112	threshold of each ear. As the criteria for normal hearing, an average air conduction value of 15
113	dB HL or less was taken (Olusanya et al., 2019). High-frequency audiometry was done for the
114	frequencies from 9 kHz to 16 kHz. The six frequency average of 9 kHz, 10 kHz, 11.2 kHz, 12.5
115	kHz, 14 kHz, and 16 kHz was taken to determine the threshold.
116	Biologic Navigator Pro equipment was used to record auditory brainstem response (ABR) for all
117	the participants. The recording was done in the soundproof room following the ANSI guidelines
118	(Frank., 1997). The participants were instructed about the procedure and aim of the test before
119	starting the recording. The participants were asked to sit in the reclining chair and ensure they
120	were comfortable enough to begin the test. The participants were made ready for the test with
121	proper cleaning, and they were instructed to sleep and relax during the entire testing to minimize
122	the artifact and stabilize the electroencephalogram.
123	The single-channel recording was done in all the participants with vertical electrode montage.
124	Test ear (A1 or A2) was used as the inverting electrode site (-), the Upper forehead (Fpz) was as
125	the non-inverting electrode (+), and the contralateral ear of the test ear was used as the common
126	ground electrode site using the 10-20 international electrode site classification (Homan et al.,
127	1987). Cup electrodes were used for recording all the participants. Electrode Impedance of $3k\Omega$
128	and inter-electrode impedance of $1k\Omega$ was maintained during the entire recording for all the
129	participants. To deliver the stimulus, Radio ear Insert-3A was used as the transducer. The click
130	stimulus of 100-microsecond duration was used as the stimulus at the intensity of 90 dB SPL.
131	The recording was done at two different rates, 11.1/s, and 90.1/s, with rarefaction polarity.
132	The acquisition parameters used were a filter setting of 100Hz to 1500Hz, amplification of
133	$1,00,000$ times, a time window of 10 milliseconds, and artifact rejection of $23.6\mu V$ (Hurley.,
134	2012). The averages of 1500 were taken, and consistency was maintained for all the participants
135	included in the study. During the entire testing procedure, it was made sure that the
136	electroencephalogram (EEG) was within the standard limit. The recording was done in all the
137	participants with replication for reproducibility. The three experienced audiologists identified the
138	peaks following the criteria established in the literature, with visualization of three sequences of
139	the peaks as I-III-V using Bio-logic Auditory Evoked Potentials (Ver 7.2.1) software. The
140	absolute latency, interpeak latency, inter-rate latency difference, and amplitude of ABR peak, i.e.
141	I, III, and V were calculated and analyzed between the misophonia and control group.

The IBM SPSS program, version 25.0, was used for the data analysis. The Sapiro-Wilk test was carried out to determine the normality. As the data followed a normal distribution, a parametric one-way ANOVA test was conducted to find the significant differences between the misophonia and control groups. The dependent variables were the latency and amplitude of all the peaks, and the independent variable was the severity of misophonia. The criteria for statistical significance was set at a p-value of less than 0.05 with a 95% confidence interval

3.0 RESULTS

3.1 Misophonia Severity

We found that 10 participants had moderate to severe misophonia with scores ranging from 21 to 30, 5 participants had severe to extreme misophonia with a score ranging from 31 to 40, and 15 participants had mild misophonia with a score ranging from 11-20. Altogether, 30 participants were included in the misophonia group. All 15 participants who were included as the control group had a score of zero on the revised RAMISO-S scale. We did not have enough data for the misophonia group to form the three groups; hence, we divided participants into two groups, one mild misophonia group and another moderate-severe misophonia group. The mild misophonia group had a score of 15.93 (SD=2.89), and the moderate-severe misophonia group (N=15) with a mean score of 25.86 (SD=4.98). All the participants included in the study had misophonia for 4.9 years with a variation from 3 to 8 years (Mean=4.93, SD=1.52).

3.2. Audiological evaluation

The physical examination showed a normal appearance of the external and middle ear in all the participants. All the participants had normal health conditions with normal hearing. The result did not show a statistically significant difference in the air conduction threshold between the study and control groups with (F (2.42) = 0.587, p= 0.561) for the right ear and with (F (2.42) = 2.540, p= 0.091) for the left ear. Similarly, we did not find statistically significant differences between the study and control groups for the bone conduction threshold also, with (F (2.42) = 0.678, p= 0.66) for the right ear and with (F (2.42) = 1.540, p= 0.08) for the left ear as illustrated in **table 1.**

The results of the high-frequency audiometry showed the presence of normal hearing in the high
frequency range from 9 kHz to 16 kHz for all the participants. The ANOVA result did not show
any significant difference in the mean high-frequency average between the groups, with (F (2.42
= 3.401, p= 0.062) for the right ear and with (F (2.42) = 1.769, p= 0.183) for the left ear. Table 1
shows the audiological findings of pure-tone and high-frequency audiometry.
Insert Table 1 here
3.3 Auditory Brainstem Response (ABR) Findings
The result of the Auditory Brainstem Response (ABR) was analyzed to determine the neural
processing in the retro-cochlear pathway. During the entire recording, electrode impendence was
less than $3k\Omega$, and the inter-electrode difference was less than $1k\Omega$ (Hurley., 2012). The absolute
latency, the amplitude of all the peaks, interpeak latency, and latency difference at the different
rate was analyzed for all the participants.
Absolute Latency
Absolute latency of I, III, and Vth peaks was analyzed between the groups at two different rates:
11.1/sec and 90.1/sec. For the rate of 11.1/sec, the result of the one-way ANOVA showed no
significant differences between the group for the absolute latency values of the I peak, III peak,
and V peak for both ears (p>0.05). Similarly, for the rate of 90.1/sec, we did not find a
significant difference between the groups for the absolute latency values of the I peak, III peak,
and V peak for both ears p>0.05). The mean and standard deviation of all the peaks of ABR,
along with the one-way ANOVA result, are illustrated in Table 2 and Table 3 for the right ear
and left ear, respectively.

193

Insert Table 3 here

194

195

Amplitude of peaks

- The amplitude of all the peaks of ABR was analyzed between the groups. The rate used was
- 11.1/s. The mean value and standard deviation of all the peaks of ABR are illustrated in Figure 2
- and Figure 3, respectively.

Insert Figure 1 here

199

Insert Figure 2 here

200

- The result of the one-way ANOVA showed no significant differences in the amplitude of all the
- 202 peaks for both ears. For the amplitude of the I peak, no significant difference was found between
- 203 the groups with (F(2.42) = 1.62, p = 0.21) for the right ear and (F(2.42) = 1.52, p = 0.31) for the
- left ear. Similarly, for the amplitude of the III peak, we did not find any significant difference
- between the groups with (F (2.42) = 1.19, p= 0.32) for the right ear and (F (2.42) = 1.52, p=
- 206 0.31) for the left ear. For the Vth peak also, no significant differences were found between the
- groups with (F(2.42) = 0.31, p = 0.74) for the right ear and (F(2.42) = 0.35, p = 0.71) for the left
- 208 ear.

209

Interpeak latency (IPL)

- 210 Interpeak latency difference of I and III peaks, III and Vth peak, and I and Vth peak were
- analyzed at the rate of 11.1/s between the control group and the misophonia group. The result of
- 212 the one-way ANOVA showed no significant difference between the groups for all the interpeak
- 213 latency differences (p>0.05) as illustrated in Table 4.

Insert Table 4 here

Inter-rate latency difference

The absolute latency difference at two rates: 11.1/s and 90.1/s, was calculated by subtracting the absolute latency value at 11.1/sec from 90.1/sec. The comparison was made between the two misophonia groups and the control group. The mean value of absolute latency difference at two different rates (11.1/s and 90.1/s) was found to be less than 0.8ms suggesting no indication of retrocochlear pathology. The mean and standard deviation value for rate differences values for all the peaks along the ANOVA result is illustrated in **Table 5.** The result of the one-way ANOVA showed no significant difference in the rate difference values for all the peaks (p>0.05), as illustrated in Table 5.

Insert Table 5 here

DISCUSSION

misophonia through auditory brainstem response (ABR) testing. The ABR was performed on all the participants, and responses were analyzed between the control and misophonia groups. The absolute latency of the peaks, amplitude, interpeak latency differences, and inter-rate latency differences was calculated and analyzed among all the participants recruited in the study. The comparison of the ABR parameters of the misophonia group with the control group showed no significant differences in all the ABR parameters. These results showed the presence of normal retro-cochlear pathway processing up to the brainstem structures among individuals with misophonia.

Misophonia is a disorder that may occur alone or in association with other auditory disorders, such as tinnitus and hyperacusis (Dozier., 2015). There has been growing interest in the use of ABR among individuals with tinnitus. Various studies have shown an increase in latency and a decrease in amplitude of all the peaks of ABR among individuals with tinnitus (Sand & Saunte., 1994; Keith & Greville, 1987). As misophonia occurs in association with these auditory disorders, we hypothesized that there could be some differences in neural processing at the level of the peripheral nervous system among individuals with misophonia. However, the result of our

Our study aimed to analyze the processing of the brainstem pathway in individuals with

243	study showed no significant differences in the ABR parameters among individuals with
244	misophonia, rejecting the hypothesis. The difference in these findings among tinnitus and
245	misophonia showed a difference in pathophysiology among these disorders. However, this is the
246	first study of this kind among misophonia and needs to replicate the findings in the future, taking
247	a larger sample size.
248	Various neuroimaging investigations using functional magnetic resonance imaging (fMRI) has
249	shown abnormal processing of the various auditory cortical areas, including non-classical
250	auditory pathway among individuals with misophonia (Kumar et al., 2017; Brout et al., 2018).
251	However, no neuroimaging studies have reported abnormal processing of sub-cortical auditory
252	pathways, including brainstem areas, among individuals with misophonia. Our study also
253	supports neuroimaging studies' findingss, suggesting normal processing of retrocochlear
254	structures among individuals with misophonia. These findings from the electrophysiological and
255	neuroimaging investigation suggest retro-cochlear structure abnormalities are absent among
256	individuals with misophonia.
257	In our study, we assessed the ABR using both low and high stimulation rates. Both rate levels
258	did not show any significant difference in the ABR findings among the control and misophonia
259	groups. These results suggest the presence of normal neural synchrony up to brainstem areas
260	among individuals with misophonia. However, our study could not obtain frequency-specific
261	ABR responses as we used the click stimulus (Robier et al., 1992). Hence, there is a need to
262	carry out studies in the future using frequency-specific stimuli.
263	
264	CONCLUSION
265	Our study concludes normal processing of the retro-cochlear pathway among individuals with
266	misophonia. However, this is the first study of this kind, and further studies on the larger
267	population are needed to generalize the results. The result of our research would be the baseline
268	for all the neurologists, audiologists, and psychologists working in misophonia to understand the
269	disorder.

2/1	Limitations and future directions
272	There is a shortage of studies in the literature assessing misophonia from the audiological
273	perspective. Our study showed normal auditory brainstem responses in individuals with
274	misophonia. The result of our research using the electrophysiological measure supports the
275	findings of the functional Magnetic Resonance Imaging (fMRI) studies reported in the literature.
276	However, this is the first study of this kind from the audiological perspective. We need to
277	validate our findings in the future, taking a larger sample size. In addition, we need to carry out
278	studies in the future using frequency-specific stimuli. Furthermore, our study used the most
279	widely used questionnaire RAMISO-S for assessing misophonia and its severity, which is
280	unavailable in the native Indian language and population. There is a need to carry out the study
281	in the future, using the questionnaire standardized in the native language.
282	
283	
284	DATA AVAILABILITY STATEMENT
285	The datasets used for the current study are available from the corresponding author upon
286	reasonable request.
287	
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355	Figure legends
356	Figure 1: Representation of amplitude of all the peaks of ABR for the Right ear (N=45)
357	Figure 2: Representation of amplitude of all the peaks of ABR for the left ear (N=45)
358	

		HEARING THRESHOLD Pure tone average (dBHL)			ESHOLD	High frequency average (dB SPL)	
		Right ear		Left ear		Right	Left ear
		Air conduction	Bone Conduction	Air Conduction	Bone Conduction	ear	
CONTROL	Mean	6.08	3.5	7.75	4.8	7.47	8.36
	N	15	15	15	15	15	15
	Std. Deviation	3.07	1.7	3.69	2.2	2.57	4.16
MILD	Mean	7.75	3.2	6.08	4.5	8.33	10.72
	N	15	15	15	15	15	15
	Std. Deviation	7.58	2.5	4.60	1.5	3.34	5.16
MODERA	Mean	8.33	4	7.75	4.5	8.43	11.28
TE-	N	15	15	15	15	15	15
SEVERE	Std. Deviation	6.58	1.7	3.69	2.2	3.05	4.09
F-value		0.59	0.68	2.54	1.54	3.40	1.77
Sig.		0.56	0.66	0.09	0.08	0.06	0.18

Table 1: Result of the pure tone audiometry and high frequency audiometry between the groups for the Right ear and left ear (N=45)

ABSOLUTE LATENCY (ms)

			11.1/sec			90.1/sec	
		I	III	\mathbf{V}	I	III	\mathbf{V}
CONTROL	Mean	1.44	3.48	5.08	1.56	3.69	5.66
	N	15	15	15	15	15	15
	Std. Deviation	0.19	0.16	0.37	0.24	0.16	0.27
MILD	Mean	1.36	3.56	5.15	1.56	3.65	5.62
	N	15	15	15	15	15	15
	Std. Deviation	0.12	0.49	0.19	0.23	0.36	0.20
MODERA TE- SEVERE	Mean	1.43	3.55	5.22	1.46	3.81	5.66
	N	15	15	15	15	15	15
	Std. Deviation	0.12	0.15	0.27	0.43	0.19	0.28
F-value		1.37	0.90	0.95	0.48	1.71	0.13
Sig.		0.26	0.41	0.39	0.62	1.93	0.88

Table 2: Result of one-way ANOVA showing the absolute latency of I, III, and Vth peaks at two different rates (11.1/s and 90.1/s) for the right ear

ABSOLUTE LATENCY (ms)

			11.1/sec			90.1/sec	
		I	III	\mathbf{V}	I	III	\mathbf{V}
CONTROL	Mean	1.50	3.54	5.13	1.53	3.78	5.71
	N	15	15	15	15	15	15
	Std. Deviation	0.14	0.12	0.29	0.45	0.17	0.22
MILD	Mean	1.37	3.58	5.25	1.49	3.59	5.72
	N	15	15	15	15	15	15
	Std. Deviation	0.39	0.19	0.22	0.63	0.86	0.14
MODERA TE- SEVERE	Mean	1.47	3.61	5.31	1.66	3.95	5.79
	N	15	15	15	15	15	15
	Std. Deviation	0.08	0.16	0.29	0.14	0.23	0.26
F-value		1.06	0.66	1.62	0.55	1.31	0.69
Sig.		0.36	0.52	0.21	0.58	0.28	0.51

Table 3: Result of one-way ANOVA showing the absolute latency of I, III, and Vth peaks at two different rates (11.1/s and 90.1/s) for the left ear.

		INTER-PEAK LATENCY DIFFERENCE (ms)						
		RIGHT EAR				LEFT EAR		
		I-III	III-V	I-V	I-III	III-V	I-V	
CONTROL	Mean	2.04	1.59	3.64	2.05	1.59	3.63	
	N	15	15	15	15	15	15	
MILD	Std. Deviation	0.14	0.35	0.30	0.13	0.30	0.27	
	Mean	2.20	1.60	3.80	2.00	1.67	3.49	
	N	15	15	15	15	15	15	
	Std. Deviation	0.22	0.21	0.21	0.57	0.33	0.98	
MODERA TE- SEVERE	Mean	2.12	1.67	3.64	2.14	1.69	3.83	
	N	15	15	15	15	15	15	
	Std. Deviation	0.16	0.25	0.66	0.17	0.23	0.29	
F-value		3.18	0.36	0.71	0.65	0.57	1.19	
Sig.		0.05	0.70	0.49	0.53	0.57	0.32	

Table 4: Result of one-way ANOVA showing Interpeak latency difference between the groups for the Right ear and Left ear at the rate of 11.1/s

INTER-RATE ABSOLUTE LATENCY DIFFERENCE (90.1/S-11.1/s) **RIGHT EAR LEFT EAR** I III \mathbf{V} Ι \mathbf{V} Ш **CONTROL** 0.14 0.20 0.14 0.24 Mean 0.58 0.58 N 15 15 15 15 15 15 Std. Deviation 0.11 0.08 0.26 0.15 0.11 0.29 **MILD** Mean 0.24 0.23 0.44 0.22 0.25 0.46 N 15 15 15 15 15 15 Std. Deviation 0.23 0.16 0.22 0.15 0.17 0.23 Mean 0.16 0.28 0.44 0.19 0.34 0.49 **MODERAT E-SEVERE** N 15 15 15 15 15 15 0.14 0.16 0.30 Std. Deviation 0.17 0.15 0.13 2.03 F-value 1.69 0.74 1.45 1.17 2.09 0.25 0.49 Sig. 0.19 0.15 0.32 0.14

Table 5: Result of ANOVA showing absolute latency difference at two different rate (11.1/s and 90.1/s) between the groups for the Right ear and left ear (N=45)



