

4 **The effects of familiar voices on the level of consciousness**
5 **among comatose patients: a single-blind randomized**
6 **controlled trial**

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11 **ABSTRACT**

Background: Brain injury can reduce consciousness and the ability to respond to environmental stimulation.

Objectives: The aim of this study was to investigate the effects of familiar voices on the level of consciousness (LOC) among comatose patients with a brain injury hospitalized in the intensive care unit.

Methods: In this randomized controlled trial, sixty comatose patients with head trauma were conveniently selected from an intensive care unit of a hospital in Rasht, Iran, and randomly allocated to either a control or an intervention group. Participants in the intervention group received auditory stimulation for three consecutive days and the level of consciousness was compared in two groups. The Glasgow Coma Scale was used to assess the patients' level of consciousness. The data were analyzed through the Chi-square, the paired-samples *t*, student's *t* test, and the repeated-measures analysis of variance.

Results: A significant increase was found in the mean LOC in the intervention group after every daily auditory stimulation ($P < 0.05$). However, no significant changes were observed in the control group ($P > 0.05$). The repeated-measures analysis of variance revealed that the time and interaction of time and groups were statistically significant ($P < 0.001$).

Conclusion: Auditory stimulation with familiar voice was effective in improving levels of consciousness among comatose patients with a brain injury after three days.

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14 Keywords: Coma, Auditory stimulation, Sensory deprivation, Consciousness disorders

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17 **1. INTRODUCTION**

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19 Brain injury (BI) is one of the most common types of trauma (1). Annually, around ten million
20 people experience BI worldwide, of whom five million are from the United States (2). In Iran,
21 BI is the second cause of death (3).

22 BI is mostly associated with loss of consciousness and coma. Coma, in turn, is the most
23 common cause of hospitalization in intensive care unit (ICU) (4), disabilities, and death (5-8)
24 following accidents. Sensory deprivation is one of the most common aftermaths of coma and
25 hospitalization in ICU. It considerably slows recovery (9). Therefore, strategies are needed to
26 provide comatose patients in ICU with sensory stimulation in order to prevent sensory
27 deprivation.

28 Sensory stimulation is a therapeutic method which stimulates the reticular activating system
29 in the brain and facilitates the reorganization of brain activities through creating new neural
30 links (10). Auditory stimulation is one of the sensory stimuli which can be provided to
31 patients in ICU by their family members or nurses (11).

32 Several studies supported the idea and the practice of regular and organized sensory
33 stimulation for comatose patients; however, some of them reported contradictory results (12-
34 15). For instance, a study showed that familiar sensory stimulation had no significant effects
35 on level of consciousness (LOC) (16), while two other studies reported that music therapy
36 calm comatose patients (12) and direct and indirect auditory stimulation may increase their
37 LOC (14). Thus, while sensory stimulation may potentially accelerate brain plasticity,
38 controversies exist over its effectiveness. Therefore, the present study was designed and
39 conducted to produce clearer evidence regarding the effects of auditory stimulation on
40 patient outcomes.

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42 **2. MATERIAL AND METHODS**

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44 **2.1 Design and participants**

45 As a single-blind randomized controlled trial, this study was carried out on patients with head
46 trauma admitted to the ICU of Poursina Trauma Hospital, Rasht, Iran. During the three-
47 month period of the study, i.e. from 14 July to 19 October, 2014, sixty eligible patients were
48 conveniently selected. Eligibility criteria were head trauma of any cause, comatose state with
49 a Glasgow Coma Scale (GCS) score of 3–8 for 72 hours (as determined by a neurologist),
50 an age of over sixteen, an endotracheal or tracheostomy tube in place, stable hemodynamic
51 status (characterized by a blood pressure of 90 to 160 mm Hg(17), a heart rate of 60–100
52 beats per minute, a respiratory rate of 12–24 per minute, a body temperature of 35.5–38°C),
53 and no history of previous head trauma, brain pathology, convulsion, hearing loss, cardiac
54 arrest, skull fracture, intracranial hemorrhage, and surgery on the temporal lobe of the brain.
55 Exclusion criteria were patient death or hospital discharge during the study and a sudden
56 significant change in hemodynamic status. During the sampling period, 83 patients with head
57 trauma were admitted to the study setting. The legal guardians of seven patients did not
58 consent for participation, seven patients experienced death or were discharged from ICU
59 during the study, and nine had unstable hemodynamic status. Thus, the remaining sixty
60 patients were included (Figure 1).

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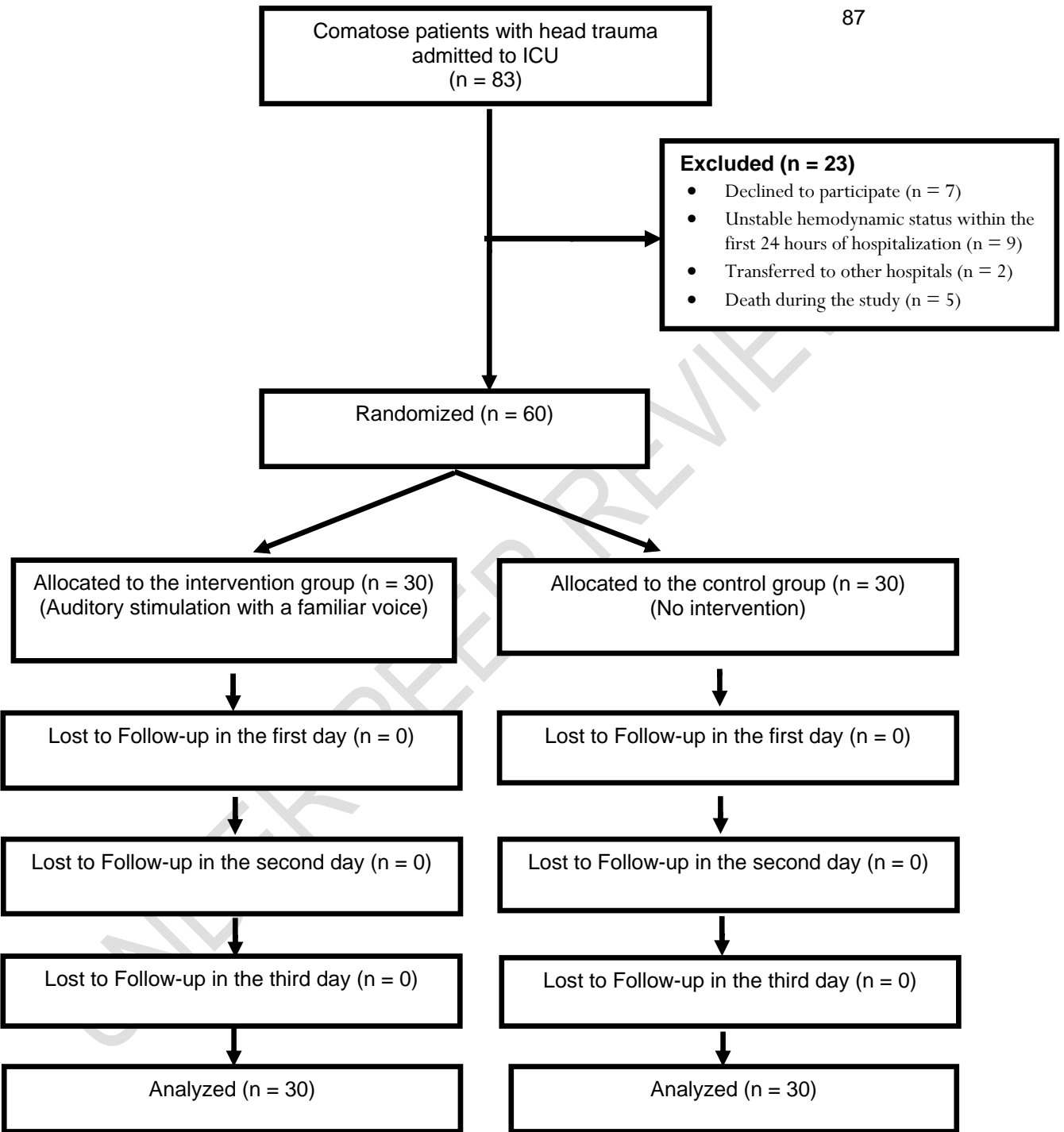


Figure 1. The CONSORT flow diagram of the study

138 Based on the findings of a previous study (13) and with a type I error of 0.01, a type II error
139 of 0.2, a μ_1 of 7, a μ_2 of 6.2, an S_1 of 0.84, an S_2 of 0.76, and a d of 0.8, sample size was
140 estimated as thirty patients per group based on the following formula.

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$$n = \frac{(Z_{1-\alpha/2} + Z_{1-\beta})^2 (S_1^2 + S_2^2)}{(\mu_1 - \mu_2)^2}$$

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145 The selected sixty participants were randomly and equally allocated to either a control or an
146 intervention group through block randomization (11). Sampling conducted based on random
147 block process by computer. As the sample size was calculated 60 patients, we used 15
148 quadruple blocks (with regard to the two existent study groups) and with concealment, 30
149 patients were allocated to intervention group and 30 individuals to control group.

151 **2.2 Data collection**

152 A four-part instrument was used for data collection. The first part included items on age,
153 gender, marital status, education level, and history of serious illnesses in the past. This part
154 was completed through interviewing participants' family members. The second part included
155 items on participants' clinical characteristics such as the cause of coma, intracranial
156 hemorrhage according to the computed tomography scan findings, surgery for intracranial
157 hematoma management, duration of coma, the need for mechanical ventilation, and
158 medications. The third part contained items on hemodynamic status, namely mean arterial
159 pressure, heart rate, respiratory rate, and body temperature. Data on mean arterial pressure,
160 heart rate, and respiratory rate were obtained from a bedside monitoring device. The
161 monitoring device was also calibrated before measurements. Blood pressure was measured
162 from the right hand through a non-invasive method while the head of bed was elevated by
163 thirty degrees. Body temperature was measured using a mercury-in-glass thermometer. The
164 fourth part was the fifteen-item GCS. The content validity of the first three parts of the
165 instrument was confirmed by ten nursing and medical faculty members.

168 **2.3 Intervention**

169 The study intervention was auditory stimulation through familiar voices. Accordingly, the
170 family of each patient in the intervention group was asked to introduce one of its members
171 who had the closest relationships with the patient. Then, the family members were trained
172 about how to record a ten-minute voice message. The first part of the message was included
173 the information about time and place (thirty seconds) and the accident which had lead to
174 head trauma (thirty seconds).

175 In the second part that lasted four minutes, they talked about shared sweet memories. In the
176 third part, they spoke promising and encouraging words about the patient's recovery and
177 future subjects (17) (five minutes). This message was recorded in the visitation room of the
178 ICU in the first 24 hours after recruitment to the study and using a voice recorder (LD-73,
179 Lander electronics).The recorded audio files were played for the intended patient in three
180 consecutive days in the afternoon, before the patient's visit time (13).The LOC was
181 assessed using GCS, both five minutes before and five minutes after each auditory
182 stimulation session Moreover, hemodynamic parameters were measured both two minutes
183 before and two minutes after the intervention (17).

184 Data were collected by the first author who was aware of the allocation sequence. Patients
185 in the control group received no auditory stimulation; but their LOC and hemodynamic
186 parameters were assessed in the same time points as their counterparts in the intervention
187 group.

189 **2.4 Ethical considerations**

190 At the time of sampling, the aim of the study was explained to participants' family members
191 and their informed consent was obtained. They were assured of the confidentiality of their
192 patients' information as well as the voluntariness of participation in and withdrawal from the
193 study. Moreover, we did our best to protect participants' rights according to the Declaration

194 of Helsinki. The study was approved by the Ethics Committee of Guilan University of Medical
 195 Sciences, Rasht, Iran (code: REC.9161.2930162909). It was also registered in the Iranian
 196 Registry of Clinical Trials (code: IRCT2014051517693N1).

198 **2.5 Data analysis**

199 The data were analyzed using the SPSS software v. 16.0 (SPSS Inc., Chicago, IL, USA).
 200 The Chi-square test was used for between-group comparisons in terms of nominal and
 201 ordinal variables such as gender, age, marital status, educational level, mechanism of head
 202 trauma, brain tissue injury, and the need for surgery. Moreover, the t-test was used for
 203 between-group comparisons in terms of continuous variables such as LOC. The paired-
 204 sample *t* test was also used for within-group comparisons in terms of LOC, while the
 205 repeated-measures analysis of variance was conducted to compare LOC in both groups
 206 across the three days of the study. The level of statistical significance was set at less than
 207 .05

209 **3. RESULTS**

210 Most participants were male (76.6%) and married (61.6%). Age mean in the intervention and
 211 the control groups were 35.16 ± 14.1 and 38.13 ± 13.89 , respectively. **Before intervention,**
 212 no statistically significant differences were found between the groups in terms of the baseline
 213 LOC, **demographic and clinical characteristics**, and hemodynamic parameters (Table 1).
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Table 1. Between-group comparisons in terms of participants' demographic and clinical characteristics

Characteristics	Group	Intervention N (%) or Mean±SD	Control N (%) or Mean±SD	P value
Age	16–25	10 (33.3)	5 (16.7)	.807*
	26–35	6 (20)	10 (33.3)	
	36–45	6 (20)	3 (10)	
	46–55	6 (20)	8 (26.7)	
	56–65	1 (3.3)	3 (10)	
	> 65	1 (3.3)	1 (3.3)	
Gender	Male	23 (76.7)	23 (76.7)	.619*
	Female	7 (23.3)	7 (23.3)	
Marital status	Single	12 (40)	10 (33.3)	.49*
	Married	17 (56.7)	20 (66.7)	
	Widowed	1 (3.3)	0 (0)	
Level of Education	Illiterate	5 (16.7)	3 (10)	.141*
	Below diploma	2 (6.7)	8 (26.7)	
	Diploma	12 (40)	7 (23.3)	
	University	11 (36.6)	12 (40)	
Cause of damage	Car accident	15 (50)	16 (53.2)	.508*
	Motorcycle accident	11 (36.7)	7 (23.4)	
	Other	4 (13.3)	7 (23.4)	
LOC (GCS score)		6.1±1.26	5.93±1.33	.658**
Duration of Coma (Hours)		29.76±4.7	32.56±6.72	.102**
Brain tissue injury	Yes	30 (100)	30 (100)	.145*
	No	0 (0)	0 (0)	
Undergoing surgery	Yes	15 (50)	17 (56.7)	.605*
	No	15 (50)	13 (43.3)	

* Chi-square test

** Independent t-test

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Within-group comparisons in the intervention group in each day showed that posttest value of the LOC was significantly greater than the pretest value ($P < .05$). Though; the patients were still in coma. No significant changes were observed in the control group in this regard ($P > .05$; Table 2).

Table 2. patients' daily LOC scores at different times

Group	Group						P_b	P_c
	Intervention (Mean \pm SD)			Control (Mean \pm SD)				
Day	5 minutes before	5 minutes after	P_a	5 minutes before	5 minutes after	P_a		
First	5.43 \pm 1.1	5.73 \pm 1.33	0.005	5.73 \pm 1.14	5.76 \pm 1.13	< .326	< .305	< .999
Second	5.76 \pm 1.19	6.33 \pm 1.39	0.001	5.76 \pm 1.19	5.8 \pm 1.18	< .326	< .908	< .097
Third	6.4 \pm 1.32	6.93 \pm 1.59	0.001	5.96 \pm 1.42	6.03 \pm 1.42	< .161	< .224	< .081

a Paired- t -test for the comparison of LOC before and after the intervention

b Independent-sample t -test for the comparison of LOC in the two groups before the intervention

c Independent-sample t -test for the comparison of LOC in the two groups after the intervention

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The results of the repeated-measures analysis of variance illustrated significant increase in the posttest mean scores of LOC in intervention group across the three measurement time points ($P < .001$). However, no significant difference was observed in the control group respecting the variations of the posttest mean scores of LOC over time. There was significant difference in the interaction of time and group ($P < .001$) (Table3).

Table 3: A repeated measures ANOVA to compare mean scores of Glasgow Coma Scale in organized auditory stimulation and control group

Sum of variables	Sum of square	df	Mean square	F	Significant
<u>Within groups</u>					
Time	16.233	.766	9.194	33.075	< .001
Time x groups	6.633	.766	3.757	13.515	< .001
Between groups	11.250	1	11.250	2.226	< .141

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No significant differences were observed between the two groups in terms of hemodynamic parameters, namely mean arterial pressure, heart rate, respiratory rate, and body temperature ($P > .05$).

4. DISCUSSION

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Findings showed no significant difference between the groups in terms of LOC variations across the three measurement time points. However, there was a significant increase in LOC from the first to the third day in the intervention group. The interaction of time and group was significant that shows LOC of patients in two groups at different stages of the time after the intervention has changed differently.

248 Consistent with our findings, an earlier study reported significant increase in LOC after
249 auditory stimulation via familiar voices in intervention group (18). The findings of another
250 study reported significant difference in LOC in the study groups after a ten-day familiar
251 sensory stimulation (13). Longer duration of intervention in that study compared to the three-
252 day intervention of the present study may account for this discrepancy between these two
253 studies. Moreover, another study into the comparison of the effects of a three-day auditory
254 stimulation intervention reported improvements in patients' LOC(13). The significant effects
255 of sensory stimulation on LOC can be attributed to the high prevalence of sensory
256 deprivation among patients in ICU as well as the positive effects of sensory stimulation on
257 the reticular activating system.

258 However, it remained unknown whether familiar voice or auditory stimulation accounted for
259 LOC improvements. Considering another group with another type of auditory stimulation
260 could answer this question. Salmani et al., (2017) conducted a study into the effects of
261 affective sensory stimulation including auditory stimulation in comatose patients during the
262 first seven days of their hospitalization. The results of the study showed significant
263 improvements in LOC in the intervention group and no significant changes in the control and
264 the placebo groups (19).

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266 The findings indicated no significant difference between the intervention and the control
267 groups in terms of participants' hemodynamic parameters. Puggina et al.,(2011) showed a
268 significant increase in the hemodynamic responses in the auditory stimulation group(20).
269 Inconsistency in the results could be due to the type of auditory stimulus and different
270 sounds that can have different effects on patient. Also it may be said that the patients in the
271 present study were in a more critical condition than the patients in other studies. In the other
272 hand the differences can be due to type of medications in these patients.

273 Another finding of the present study was that the study intervention had no adverse effects
274 on participants' brain activities. Similarly, two previous studies reported that due to its non-
275 invasiveness, auditory stimulation can improve brain activities without exerting significant
276 side effects (18, 21).

277 Among the limitations of the present study were our uncertainty about the patients' favorite
278 family members as well as the short course of the study intervention. Moreover, GCS is a
279 general LOC assessment tool (22) which is not sensitive enough to the small changes in
280 LOC. The impossibility of performing the study using a double-blind design as well as the
281 differences in participants' medical treatment regimens might also have affected the study
282 results. Future studies are recommended to use double-blind designs and provide auditory
283 stimulation with familiar voices for longer periods of time and with more than one auditory
284 stimulation session per day.

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286 **5. CONCLUSION**

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288 This study indicates that auditory stimulation with the familiar voices of patients' family
289 members may improve LOC among patients with head trauma after three days. Thus, this
290 technique can be used to improve the LOC of these patients during their ICU stay. Of
291 course, longer auditory stimulation with familiar voices may produce more significant effects
292 on the LOC.

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294 **CONSENT AND ETHICAL APPROVAL**

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296 This study was conducted after receiving the written approval of the Ethics Committee of
297 Guilan University of Medical Sciences with ethics code (No: REC.9161.2930162909). Before
298 starting the sampling, we explained to the participants in terms of the objectives of the
299 research and the data collection process. Also, written informed consent was obtained from
300 all of samples.

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COMPETING INTERESTS

The authors have no conflict of interest.

ETHICAL APPROVAL

This study was conducted after obtaining the approval of the Ethics Committee of deputy of research and technology of Guilan University of Medical Sciences with Ethical code number REC.9161.2930162909) and the Iranian Center for Clinical Practice ID(code: IRCT2014051517693N1). Before performing the sampling, the participants received descriptions in terms of the aims of the study, the method of study and their rights and their expectations at each stage of the research, and in case of willingness they sign the written inform consent for participation in the study.

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