Assessment of the Quality of Life in Patients with Epilepsy in the Populous Region of Baku

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Abstract: An article establishes assessment of quality of life, severity of seizures and emotional condition 117 patients with epilepsy, over the age of 18 (35.13±1.19) with different form epilepsy what started from 0 to 69 years (14.22±1.26) in the Machtagа village. How showed analyzes of the survey results, the patients sub clinic 62 (52.9%) and clinic 29 (24.8%) pronounced measure depression and anxiety from scale Ziqmond and high measure severity of seizures from national hospital seizure severity scale (NHS3) (20-27 scores by 54 (46.1%), (3-9 scores by 40 (34.2%) patients, adversely affecting to their quality of life, assessment by questionnaire «QOLIE -10» (27.1±0.3). Social status research revealed a high patients unemployment rate (82 (70.1%) p=0.911). Between 70 (59.8%) disability patients most had secondary group (p=0.938). The number of lower secondary education and uneducated patient prevailed among native (28 (36.8%)/7 (9.2%)) and nonnative (11 (26.8%)/4 (9.8%)) village population (p=0.547). In population research 65 (55.6%) patients were married, 52 (44.4%) patients were alone (p=0.386). During the study it was revealed influence of ethnocultural characteristics patients to measure quality of life. A form of expression of religious beliefs, which, in the form of a clear implementation of the established religious rules, had a positive effect on the quality of life of patients from the hunhar and kechan mehelle, and mysticism in relation to religion, a form of upbringing in a family without encouraging intellectual development, and even prohibitions on obtaining an education, contributed to the formation passive personality with disabilities among patients from seidler mehelle, which affected their QOL negatively.

Keywords: Epilepsy, The Quality of Life, National Hospital Seizure Severity Scale, Ethnocultural Features

1. Introduction

Epilepsy is one of the most common diseases of the nervous system in children and adults. The incidence of this disease ranges from 1.5 to 18, and in some developing countries it exceeds 30 cases per 1000 population [1-6]. The incidence rate of epilepsy is growing, according to the latest literature data (Beghi E, 2020) (61.4 / 100000), which ranges from 139.0 / 100000-48.9 / 100000.

Today, the World Health Organization (WHO) regards it as one of the most important public health problems. Everyone can get epilepsy, regardless of age, gender, or race. Suffering from this ailment, it is not uncommon to face various kinds of discrimination. They are observed in getting an education, getting a job, getting married, as well as in the family and in the close environment. In addition to the social consequences, epilepsy has enormous economic costs. A few works are devoted to the study of the relationship between persons belonging to different races and ethnic groups, socio-economic formations and epilepsy. This aspect of epilepsy remains largely unexplored. Perhaps scientific research in this direction will be able to point out the undoubted role of these factors in the structure of the incidence and prevalence of this dangerous disease.

More than 60,000 people suffer from this disease in Azerbaijan to. A series of works have been devoted to the study of epilepsy in the republic over the past 20 years. Studied its epidemiology and clinic as a whole, and in its various regions [7, 8]. As you know, the problems of people with epilepsy associated with the clinical manifestations of the disease limit their desire for self-realization and social activity, thereby reducing the indicators of their quality of life [4, 9-15].

As a result, in the framework of the studies mentioned
above, the clinical and epidemiological indicators of epilepsy were studied in one of the large villages of the capital of the republic - Mashtaga, taking into account the influence of the ethnocultural characteristics of the indigenous and non-indigenous people living in this ancient settlement of Absheron on the assessment of their quality of life.

2. Method

The study of the quality of life of patients with epilepsy over the age of 18 years, living in the village of Mashtaga, in a comparative aspect of the ethnocultural characteristics of the population in the region was carried out during the period of clinical and epidemiological study of epilepsy among the population in the Mashtaga settlement of Baku city (2017-2019). The work was carried out in accordance with the protocol of the Ethics Committee No. 11 of the Azerbaijan Medical University (AMU) dated December 29, 2019 and the guidelines for conducting epidemiological studies prepared by the Commission on Epidemiology and Forecast of the population in the Mashtaga settlement of Baku city (2017-2019). The work was carried out in accordance with the protocol of the Ethics Committee No. 11 of the Azerbaijan Medical University (AMU) dated December 29, 2019 and the guidelines for conducting epidemiological studies prepared by the Commission on Epidemiology and Forecast of the International Antiepileptic League (1993) [16]. A cohort pro-and retrospective study of the epidemiology of epilepsy, taking into account the etiological and social aspects, was carried out on the basis of data on visits to the outpatient clinic in the village of Mashtaga, Sabunchu district, for the period from 2016 to 2019. Cases of active epilepsy were recorded on the results of yard rounds, a continuous study of medical outpatient records of the combined children's and adult polyclinic department of the CCH No. 7, call records of the local ambulance station No. 14. The degree of depression (D) and anxiety (A) in patients with epilepsy was determined using the Ziqmond scale, the severity of seizures was determined by the NHS3 scale, according to which the total score was determined in the range from 1 to 27. The QOLIE-10 questionnaire was used to assess the quality of life, in which each answer was graded on a scale of 1-5, with a higher score reflecting a lower quality of life, with a final score of 0-50. The age, sex of patients, their neurological symptoms, onset and causes of the disease were determined, the nature of drug therapy was studied at the beginning of the study and after optimization.

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2. Results and Discussion

During the study period (2016-2019), 45000-46787 residents lived in the village (20371-24634 men and 24625-22153 women). The indigenous population in 2017 was 37505 people, non-indigenous people who moved from different regions of Azerbaijan - 8295 people. Among them, there were 5261 people under 18 years old (2937 boys, 2324 girls), over 18 years old - 40539 people (20763 men, 19,776 women). The entire population is represented by Azerbaijani.

Representatives of the indigenous population lived in the village compactly in three quarters (mehelle): Honhaar (descendants of the Hunnic Turkish), Kechan (descendants of the Massagets), Seidler (descendants of the Arabs), which for many centuries have passed from generation to generation, the established order of behavior, customs, faith, moral values, mythological ideas [17, 18]. Marriages between representatives of these quarters are not often observed to this day.

In the studied population of 197 patients with epilepsy (125 (63.5%) from indigenous, 72 (36.5%) from non-indigenous inhabitants), 117 persons were identified at the age from 18 to 75 years. The average age of the patients was 35.13±1.19. The average age at the onset of various forms of epilepsy (from 0 to 60 years) was 14.22±1.26. Among the identified patients, 76 (65.0%) individuals were indigenous (33 (28.2%) from Honhaar, 34 (29.1%) from Kechan, and 9 (7.7%) from Seidler mehelle), and 41 (35.0%) patients were non-indigenous residents of the village. Idiopathic epilepsy was more detected in patients from Honhaar (25 (60.9%)) and Seidler mehelle (6 (14.6%)). Symptomatic prevailed in patients from Kechan mehelle (29 (38.1%) and non-indigenous people (36 (47.4%)), (p<0.05).

The severity of seizures on the NHS3 scale in patients from Kechan mehelle was assessed significantly higher (27 points) than in non-indigenous people (20 points). This may be due to the high frequency of occurrence among them of severe consequences of perinatal brain pathology (birth trauma, hydrocephalus, infantile cerebral palsy) and craniocerebral trauma, which, apparently, is associated with untimely visits to the antenatal clinic during pregnancy.

Among patients from Honhaar mehelle, persons with moderate severity of seizures were observed relatively more often (13 (39.4%); 5 points) than from Kechan (10 (29.4%); 9 points) and Seidler mehelle (3 (33.3%); 9 points), but it was rated lower. Perhaps this is due to the desire to hide the disease, which is characteristic in the ethnic aspect for the indigenous population, as well as to the unauthorized cancellation of drug treatment, which was more often observed among patients from Seidler mehelle, and to the delayed treatment due to late detection of the disease.
According to the literature, absences, as one of the forms of manifestation of idiopathic epilepsy, can often be unnoticed by the patient's relatives until they are complicated by generalized seizures [5, 19]. Regarding cases of concealment of the disease, it can be noted that a similar phenomenon occurred in other countries of the world, for example, in most countries of Asia, parents hide the disease in their daughter, in order to preserve the opportunity to marry. In a study conducted in the southern Indian city of Kerala, 39% of citizens surveyed called epilepsy a hindrance to marriage, and 55% of patients hid their diagnosis from their spouse. This was also the case in the populous province of the Republic of the Congo, in Lubumbashi, where epilepsy also posed a threat to women due to the likelihood of losing the chance to marry [20-22]. Severe seizures with a change in consciousness, accompanied by inappropriate behavior of patients, caused bewilderment on the part of others, which negatively affected the psych emotional status of patients. As is known from the literature, severe seizures play an important role in the maladjustment of patients, which significantly affects QoL [7, 9-11, 15, 23, 24]. As can be seen from diagram 2, the psych emotional status, as determined by the Zqmond scale, was assessed low in all patients (8.91±0.17; m=5, M=13), and in non-indigenous patients, it was statistically significantly lower than in indigenous patients (p=0.046). Among ethnic subpopulations, this indicator in patients from Kechan mehelle (9.26±0.33) was lower than from Honhaar (8.39±0.30) and Seidler mehelle (8.78±0.66). In the study, the majority of patients were dominated by subclinical (62 (52.9%) cases) and clinical forms of depression and anxiety (29 (24.8%) cases). In patients with subclinical (8-10 points) pronounced signs of anxiety and depression both from indigenous (46 (52.6%)) (21 (51.5%) from Honhaar, 19 (52.9%) from Kechan, 6 (55%) from Seidler mehelle) and 16 of non-indigenous residents (53.7%), no significant difference was found in the percentage. Signs of anxiety and depression, expressed clinically (above 10 points) in non-indigenous patients (29.3%) 16 individuals), prevailed over indigenous patients (22.4%) 13 individuals), (4 (12.1%) from Honhaar, 8 (32.4%) from Kechan, 1 (11.1%) from Seidler mehelle).

There were fewer patients with no pronounced symptoms of depression and anxiety (0-7 points), 26 (22.2%) cases. Among them, 9 (17.1%) were non-indigenous and 17 (25%) were indigenous (8 (36.4%) Honhaar, 7 (14.7%) Kechan, 2 (22.2%) Seidler mehelle). In addition to the clinical manifestations of the disease (frequency and severity of seizures, the form of epilepsy) and psych emotional status, the quality of life of patients with epilepsy is also influenced by their social status [4, 9, 10, 12-15].

As follows from Table 1, with study of the social status of patients, the number of unemployed persons, both indigenous (53 (69.7%)) and non-indigenous (29 (70.7%)) was significantly higher than that of workers (laborers, drivers) (23 (30.3%) of the indigenous, 12 (29.3%) non-indigenous people). Perhaps, due to material motivation, working patients rated their QoL better than unemployed ones. Among 70 (59.8%) persons with disabilities, the majority were patients with symptomatic epilepsy from Kechan mehelle 23 (67.6%) and from non-indigenous residents 26 (63.4%). The number of patients receiving disability from Honhaar and Seidler mehelle was comparatively smaller (14 (42.4%) (p <0.05), but due to late detection and complicated course of the disease, they all received the second group of disability. Persons with disabilities assessed QoL positively, which, apparently, is associated with a certain material support. The level of education in the studied 117 patients was as follows: patients with incomplete secondary education accounted for 33.3% (39 people), and 9.4% (11 people) were generally uneducated. The reason for the low education was the severe course of the disease and the prohibitions by the parents from the indigenous inhabitants, more often from the Seidler mehelle, to attend school. Patients who received education rated QoL lower than those who were uneducated. A conscious attitude towards their disease, especially of patients from the Honhaar mehelle, where the number of educated people prevailed, led to self-stigmatization.

Patients, trying to avoid potential discrimination, independently limited social activity and ignored their own social needs, which subsequently negatively affected QoL. According to many authors, self-stigmatization in epilepsy negatively affects the QoL of patients [4, 12, 17]. A study of the marital status of patients revealed that the number of married persons (65 (55.6%) outnumbered single persons (52 (44.4%)). Patients who started a family rated QoL higher than non-family patients. Feelings of compassion and support from family members increased confidence and decreased fear of seizures, which had a positive impact on QoL. In the work of Japanese researchers, carried out in 2015, it was shown that the support of relatives improves the mental health of an individual and his quality of life [25].

Table 1. Social Status of Epilepsy Patients.

| Social status of patients | Indigenous people | Non-indigenous people | Altogether: № 117 |
|---------------------------|-------------------|-----------------------|-------------------|
|                           | Honhaar № 33      | Ketchan № 34          | Seidler No 9      |
|                           | 11 (33.3%)        | 11 (32.4%)            | 1 (11.1%)         |
| Working                   | 22 (66.7%)        | 23 (67.6%)            | 8 (88.9%)         |
| Non-employed              | 2 (4.9%)          | 53 (69.7%)            | 29 (70.7%)        |
| X2, P                     | Χ2=1.782; p=0.410 | Χ2=0.013; p=0.911     |                   |
| Disabled                  | 14 (42.4%)        | 23 (67.6%)            | 7 (77.8%)         |
| I                         | 14 (42.4%)        | 4 (11.8%)             | 2 (22.2%)         |
| II                        | 3 (8.8%)          | 4 (11.8%)             | 1 (11.1%)         |
| III                       | 4 (5.3%)          | 4 (5.3%)              | 3 (8.8%)          |

Seidler mehelle was comparatively smaller (14 (42.4%)) (p <0.05), but due to late detection and complicated course of the disease, they all received the second group of disability. Persons with disabilities assessed QoL positively, which, apparently, is associated with a certain material support. The level of education in the studied 117 patients was as follows: patients with incomplete secondary education accounted for 33.3% (39 people), and 9.4% (11 people) were generally uneducated. The reason for the low education was the severe course of the disease and the prohibitions by the parents from the indigenous inhabitants, more often from the Seidler mehelle, to attend school. Patients who received education rated QoL lower than those who were uneducated. A conscious attitude towards their disease, especially of patients from the Honhaar mehelle, where the number of educated people prevailed, led to self-stigmatization.

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During the study, some ethnic characteristics of the indigenous and non-indigenous population were studied. According to the literature, out of 318 clinical studies in recent years concerning the ethnic characteristics of populations with respect to epilepsy, 6.6% of the works showed the racial identity of the study participants, and only 1.9% attempted to analyze ethnic characteristics [12, 13]. In the studied population, epilepsy sufferers were highly religious, while believers among the sick from the indigenous population prevailed over non-indigenous ones. In ethnic subpopulations, patients differed in the form of expression of religious beliefs. Patients from Honhaar and Kechan mehelle performed religious rituals in the form of strict adherence to religious beliefs. Patients from Honhaar and Kechan mehelle differed in the form of expression of the studied population, epilepsy sufferers were highly in the religious, while believers among the sick from the indigenous subpopulations, patients differed in the form of expression of the QoL. There is an opinion that the treatment of some diseases is possible through the recitation of the verses of the Qur'an [26-28]. At the same time, the religious bias of the sick from the Seidler mehelle, set them up on the wrong path of getting rid of the disease only with the help of prayers. Using prayers to combat their fear of subsequent seizures, they refused medication, which increased the severity of the seizures, complicated the course of the disease and contributed to the deterioration of QoL. It is known that even in ancient times, epilepsy was associated with the introduction into the body of the spirit, the devil, was considered as a result of divine destiny and was healed with spells to no avail [29].

### Table 2. Results of testing of epilepsy patients questionnaire QOLIE-10.

| Points | Indigenous people (meshelles) No. 76 | Non-indigenous people No. 41 | Altogether: No. 117 | p1 | p2 |
|--------|------------------------------------|-----------------------------|---------------------|----|----|
| A-energy | 4.58±0.71 | 4.47±0.11 | 4.33±0.24 | 4.50±0.08 | 4.46±0.09 | 4.49±0.06 | 0.773 |
| In-emotions | 3.30±0.11 | 3.41±0.09 | 3.67±0.24 | 3.39±0.59 | 3.32±0.08 | 3.33±0.05 | 0.115 |
| C-activity (driving) | 3.79±0.14 | 3.32±0.21 | 3.00±0.44 | 3.49±0.12 | 3.41±0.16 | 3.46±0.10 | 0.730 | 0.077 |
| D - memory | 4.03±0.03 | 4.06±0.04 | 3.67±0.33 | 4.00±0.05 | 4.10±0.05 | 4.03±0.03 | 0.174 | 0.025 |
| E-influence AED | 3.85±0.12 | 3.62±0.13 | 3.33±0.33 | 3.68±0.09 | 3.90±0.13 | 3.76±0.07 | 0.157 |
| F-waiting for seizures | 4.15±0.13 | 3.71±0.14 | 4.33±0.29 | 3.97±0.10 | 3.95±0.12 | 3.97±0.07 | 0.887 | 0.033 |
| G-QoL, subjective | 4.06±0.12 | 4.12±0.12 | 3.89±0.26 | 4.07±0.08 | 3.95±0.10 | 4.03±0.06 | 0.401 |
| QoL-objective | 27.8±0.05 | 26.7±0.05 | 26.2±1.5 | 27.1±0.3 | 27.0±0.4 | 27.1±0.3 | 0.852 |

Note: the statistical significance of the differences between the indicators:
p1 - Indigenous and non-indigenous (according to $\chi^2$ Pearson)  
p2 - Indigenous by mehelles (by $\chi^2$ Pearson)
persons II (20 (48.8%) cases) and I group (4 (9.8%) cases), and I group (4 (9.8%) cases), the body", "energy" could be influenced by the severity of being, a form of expression of religious beliefs and education, contributed to the formation of a passive personality with intellectual development and even prohibitions on obtaining an educational qualification. The desire to preserve the marriage is positive, while the concealment of the disease negatively affected the quality of life of patients from all ethnic backgrounds.

4. Conclusion

Thus, according to the objective indicators of the QOLIE-10 questionnaire, it follows that the quality of life in all patients in the population was low (27.1±0.3). The reason for the low QoL score was the high severity and frequency of seizures, the pronounced level of impairment of the psych emotional status and social adaptation of patients with epilepsy. Certain ethnocultural characteristics of patients affected the assessment of QoL. Persons both positively and negatively: a form of expression of religious beliefs, which, in the form of a clear implementation of the established religious rules, had a positive effect on the quality of life of patients from the Honhaar and Kechan mehelle, and mysticism in relation to religion, a form of upbringing in a family without encouraging intellectual development and even prohibitions on obtaining an education, contributed to the formation passive personality with disabilities among patients from Seidler mehelle, which affected their QoL negatively. The desire to preserve the marriage is positive, while the concealment of the disease negatively affected the quality of life of patients from all ethnic groups in the studied population.

Conflict of Interest Statement

The authors declare no competing interests.

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