EFFECTS OF NURSE LED EDUCATIONAL INTERVENTION ON QUALITY OF LIFE AMONG MULTIPLE SCLEROSIS PATIENTS IN LAHORE GENERAL HOSPITAL, LAHORE

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Abstract

Objective: The purpose of study is to determine the effect of nurse led educational intervention on quality of life among multiple sclerosis patients in Lahore General Hospital, Lahore

Study Design: One-group pre-post quasi-experimental study.

Place and Duration of Study: This study was conducted at Lahore General Hospital Lahore from Marc 2024 to October 2024.

Methodology: This study employed a one-group pre-post quasi-experimental design on a sample of 100 MS patients, using purposive sampling. The research included three phases: a pre-intervention phase for baseline data collection, an intervention phase consisting of 16 educational sessions aimed at improving health literacy, fatigue management, and a post-intervention phase to reassess the same variables. Data were analyzed using SPSS version 25

Results: The result of study showed that participants were primarily young adults (45% aged 18-30) and evenly split by gender (50% male, 50% female). The majority were unmarried (78%) and illiterate (49%). Post-intervention results showed significant improvements in quality of life (from median 77.5 to 111, p < 0.001). Quality of life scores significantly improved across all groups, particularly among urban participants.

Conclusion: The nurse-led educational intervention effectively enhanced quality of life among multiple sclerosis patients. These results show that educational programs for patients are necessary to improve the outcomes of

Keywords: Nurse; Educational intervention; Quality of Life; Multiple Sclerosis; Patients

Introduction

Multiple sclerosis (MS) is a chronic autoimmune disease that anxiety, and depression (5,6). significantly diminishes an individual's quality of life by Isolation is a significant challenge for individuals with multiple

due to muscle spasms, nerve pain, or other symptoms, which unable to work due to a physical or mental disability (9). issues, as the enigma of the disease's progression and the tasks (10).

potential for harm frequently induce elevated levels of stress,

disrupting the central nervous system. The illness's sclerosis. The condition's mental and physical effects may cause unpredictable nature and the extensive array of potentially individuals to avoid social situations due to their belief that incapacitating symptoms present numerous obstacles for others do not comprehend them or their reluctance to disturb individuals with the condition (1). Physical limitations, their family and friends. Tiredness, difficulty moving, and low including muscle weakness, stiffness, and fatigue, can self-esteem all contribute to their sense of disconnection and complicate daily activities and require the use of mobility aids inadequacy in social situations, which impedes their like wheelchairs or walkers. These modifications frequently participation in social activities (7,8). Additionally, the expenses undermine an individual's sense of autonomy, leading to associated with medications, medical apparatus, and housing feelings of frustration, isolation, and diminished self-esteem modifications frequently exacerbate the financial anxiety associated with multiple sclerosis. Patients and their families In addition, certain individuals with MS experience chronic pain may experience additional financial difficulties if they are

hinders their ability to engage in activities that they previously The disease significantly impacts the relationships of found enjoyable. Cognitive disorders, including amnesia, individuals, as caretakers assume additional responsibilities and difficulty concentrating, and difficulty managing problems, can patients contend with feelings of remorse or dependence. significantly impede one's ability to engage in work, fulfil daily Mental distress may result from the gradual loss of obligations, and maintain relationships with friends and family independence, which can lead to a decrease in self-esteem and (4). The emotional pain is exacerbated by physical and mental an increased dependence on others for personal care and daily

Methods

questionnaire, as cited in (11). This extensive 31-component and 17 (17.0%) were situated in towns. evaluation examines multiple dimensions of quality of life, including emotional well-being, physical functioning, and Quality of Life of Multiple Sclerosis Patients sessions covering topics related to MS related quality of life group management. Pre- and post-intervention data were analyzed using SPSS, with Wilcoxon Signed Rank and Chi-Square tests applied to compare the results, considering a p-value of less than 0.05 as statistically significant.

Results

Table 4.1. Demographic information of participants

| Study Variable | Category | Frequency (F) | Percentage (%) |
|-------------------|----------------|---------------|----------------|
| | <18-30 Years | 45 | 45.0 |
| A 000 | 31- 40 Years | 40 | 40.0 |
| Age | 41- 45 Years | 9 | 9.0 |
| | > 45 Year | 6 | 6 |
| Gender | Male | 50 | 50 |
| | Female | 50 | 50 |
| Marital Status | Married | 20 | 20 |
| | Unmarried | 78 | 78 |
| | Widow/Divorced | 2 | 2 |
| Educatio | Illiterate | 49 | 49 |
| n Level | Primary school | 51 | 51 |
| Residenc e | Rural | 27 | 27 |
| | Urban | 56 | 56 |
| | Town | 17 | 17 |

Table 4.1 provides the demographic information of the study participants, categorized by age, gender, marital status, Figure 1: Multiple Sclerosis Quality of Life Assessment education level, and residence. Among 100 enrolled multiple sclerosis patients, 45 (45.0%) were aged between 18 and 30 years, 40 (40.0%) were aged between 31 and 40 years, 9 (9.0%)

were aged between 41 and 45 years, and 6 (6.0%) were over 45 The research utilised a one-group pre-post quasi-experimental years of age. In terms of gender distribution, the cohort was framework to evaluate the effects of a nurse-led educational evenly split, with 50% male and 50% female participants. intervention on the quality of life for individuals diagnosed with Marital status revealed that 20 (20.0%) of the patients were multiple sclerosis. The educational intervention served as the married, 78 (78.0%) were unmarried, and 2 (2.0%) were independent variable, while the quality of life functioned as the widowed or divorced. Regarding educational attainment, 49 dependent variable. A cohort of 100 individuals diagnosed with (49.0%) of the subjects were illiterate, while 41 (41.0%) had multiple sclerosis was employed to collect data at Lahore completed primary school, and 10 (10.0%) had completed high General Hospital. The evaluation of quality of life was carried school. The living arrangements of the patients indicated that 27 out using the Multiple Sclerosis Quality of Life (MSQOL) (27.0%) resided in rural areas, 56 (56.0%) lived in urban areas,

social engagement. The intervention spanned four educational Table 4.2: Quality of Life among pre and post intervention

| Quality of Life | Pre Median (IQR) | Post Median (IQR) | Wilcoxo n Signed Ranks Test | p- value |
|--|------------------------|-------------------------|--------------------------------------|-------------|
| Multiple Sclerosis Quality of Life (MSQOL- 31) | 77.5(70.2 5-85) | 111(105- 114) | -8.592° | <0.00 1* |

Table 4.2 highlights the significant improvement in the quality of life for participants with multiple sclerosis following the intervention. The median score on the Multiple Sclerosis Quality of Life (MSQOL-31) scale increased from 77.5 (IQR: 70.25-85) before the intervention to 111 (IOR: 105-114) after the intervention. The Wilcoxon Signed Ranks Test result of -8.592c and a p-value of <0.001 indicate a statistically significant improvement in quality of life post-intervention, showing the positive impact of the intervention on the participants' wellbeing.

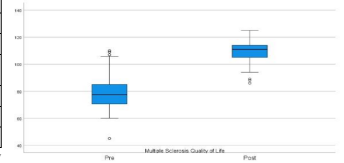


Table 4.3: Ouality of Life of pre and post intervention group according to demographic variables

| Variable | Pre Interven | Pre Intervention Quality of Life | | Post Intervention Quality of Life | |
|------------|--------------|----------------------------------|-----------|-----------------------------------|--|
| Age | Low | Average | Average | Good | |
| 18-30 Year | 21(46.7%) | 24(53.3%) | 37(82.2%) | 8(17.8%) | |
| 31-40 Year | 18(45%) | 22(55%) | 30(75%) | 10(25%) | |
| 41-45 Year | 6(66.7%) | 3(33.3%) | 7(77.8%) | 2(22.2%) | |
| > 45 Year | 5(83.3%) | 1(16.7%) | 3(50.0%) | 3(50.0%) | |
| P-Value | 0.210 | 0.210 | | 0.408 | |

| Gender | | | | |
|------------------------|-----------|-----------|-----------|-----------|
| Male | 28(56.0%) | 22(44.0%) | 37(74.0%) | 13(26.0%) |
| Female | 22(44.0%) | 28(56.0%) | 40(80.0%) | 10(20.0%) |
| P-Value | 0.230 | | 0.476 | |
| Marital Status | | | | |
| Married | 8(40.0%) | 12(60.0%) | 15(75.0%) | 5(25.0%) |
| Unmarried | 41(52.6%) | 37(47.4%) | 62(79.5%) | 16(20.5%) |
| Widow/ | 1(50.0%) | 1(50.0%) | 0(0%) | 2(100.0%) |
| Divorced | | | | |
| P-Value | 0.603 | | 0.045* | |
| Education Level | | | | |
| Illiterate | 18(36.7%) | 31(63.3%) | 41(83.7%) | 8(16.3%) |
| Primary School | 32(62.7%) | 19(37.3%) | 36(70.6%) | 15(29.4%) |
| P-Value | 0.009* | | 0.120 | |
| Residence | | | | |
| Rural | 20(74.1%) | 7(25.9%) | 15(55.6%) | 12(44.4%) |
| Urban | 20(35.7%) | 36(64.3%) | 49(87.5%) | 7(12.5%) |
| Town | 10(58.8%) | 7(41.2%) | 13(76.5%) | 4(23.5%) |
| P-Value | 0.003* | | 0.007* | |

Table 4.3 presents the quality of life outcomes in the pre- and including the Middle East and Europe, indicates that women are that marital status had a significant impact post-intervention, effective than it may have been. particularly among widowed/divorced individuals (p = 0.045). The educational intervention considerably enhanced the quality Education level played a key role in determining the quality of of life, shown by a median score of 111, an increase from 77.5 improvement.

Discussion

causes inflammatory demyelination and axonal damage in the were statistically significant in relation to marital status central nervous system. It is associated with a number of (P=0.045), education level (P=0.009), and residence (P=0.007), contributing factors (14,15).

General Hospital population impacted by multiple sclerosis interventions. (MS). Half of the people who took part were between the ages This finding is consistent with research by (25), which of 18 and 30, and another half were between the ages of 31 and highlighted that patient education improved not only health 40. The age distribution is in line with global evidence showing outcomes but also quality of life in chronic illness management. that multiple sclerosis often manifests in early adulthood, However, the degree of improvement varied by socioeconomic namely between the ages of 20 and 40 (16). Research conducted and educational status, which mirrors the present study's in Turkey supports these findings, showing that persons aged findings regarding the impact of education on MS patients' 20–40 also have the highest prevalence of MS (17,18).

Fifty percent of the responders were male and fifty percent were reported that education of patients improved the quality of life female, indicating an even gender distribution. This sharply of patients. Similarly, these findings are consistent with other contrasts with existing statistics, which typically indicate that study who reported that coping strategies have positive impact females have a higher prevalence of MS, sometimes at a ratio of on the quality of life (27). 2:1 or more (19,20). Research conducted in several regions,

post-intervention groups, categorized by various demographic more susceptible to multiple sclerosis (21). Fifty-one percent of variables. It examines differences in quality of life based on age, respondents had a high school diploma or its equivalent, whilst gender, marital status, education level, and residence. In the age forty-nine percent were entirely illiterate. Research indicates category, individuals aged 18-30 and 31-40 showed a higher that elevated educational attainment correlates with enhanced shift towards improved quality of life post-intervention, while health literacy and improved ability to manage chronic illnesses those over 45 had mixed outcomes. Gender-wise, both males (22). Patients with lower literacy skills had more challenges in and females saw a substantial improvement post-intervention, comprehending health-related materials, perhaps rendering the with females slightly outperforming males. The data also shows educational intervention implemented by nurses in this trial less

life, with illiterate individuals demonstrating more improvement before to the intervention (p < 0.001). These findings align with compared to those with primary education (p = 0.009 pre- those of (23), which indicated that educational programs for intervention). Finally, residence was a notable factor, with urban multiple sclerosis (MS) patients may significantly improve their participants showing the most significant improvement in quality of life by addressing both physical and emotional quality of life post-intervention (p = 0.003 pre-intervention and dimensions of treatment. Recent evidence from a three-year p = 0.007 post-intervention), while rural participants had lesser longitudinal study indicates that patient education enhances quality of life by alleviating psychological distress and improving functional outcomes (24).

The current study revealed that the quality of life of MS patients An autoimmune disease known as multiple sclerosis (MS) was improved after nursing intervention. These improvements suggesting that these factors may influence how well patients This study's demographic profile sheds light on the Lahore respond to health literacy and fatigue management

quality of life. These findings are consistent with (26) which

Conclusion

were young and aged between 18-30 years, with an equal gender sclerosis: Analysis of ten years of national data. Multiple distribution (50% male, 50% female). The majority were Sclerosis and Related Disorders, 66, 104019. unmarried (78%) and had a primary school education (51%). In 5. terms of residence, 56% were from urban areas, while 27% lived well-being, and sleep quality in multiple sclerosis. Journal of in rural areas. The study's findings indicate significant clinical medicine, 12(2), 716. improvements in quality of life among multiple sclerosis 6. patients following the intervention. Health literacy scores Emanuele, C., Katia, M., Caterina, F., Carmela, R., Edoardo, increased substantially, with younger age groups and unmarried S., & Giangaetano, D. A. (2022). Psychological well-being in participants showing the most progress. Quality of life improved people with multiple sclerosis: a descriptive review of the post-intervention, particularly among urban residents, illiterate effects obtained with mindfulness interventions. Neurological and widowed/divorced participants. intervention was notably effective, positively impacting 7. patients' overall well-being.

Limitations of Study

- The sample size was limited, which limit the 734. generalizability of the results to a broader population of multiple 8. sclerosis (MS) patients.
- General Hospital), which limit the applicability of findings to other settings or regions with different healthcare infrastructures 9. or patient populations.
- A large portion of the participants had low educational backgrounds, which might have influenced the outcomes. The multiple sclerosis: A cross-sectional study. Multiple Sclerosis study does not explore how the intervention would perform with and Related Disorders, 59, 103641. patients having higher levels of education.

Recommendations of study

- educational interventions tailored to the specific needs of Emotional Alterations in Patients with Multiple Sclerosis. patients with multiple sclerosis.
- Stay updated on evidence-based practices to improve health, 20(2), 1387. patient outcomes in managing multiple sclerosis.
- holistic care that addresses both the physical and psychosocial Beiske, A., & Vollmer, T. (2008). Validation of the multiple needs of multiple sclerosis patients.
- Recommendations for Patients 4.
- Engage actively in educational sessions to gain a better 12. 5. understanding of multiple sclerosis and its management.

Utilize provided resources and support systems to improve selfmanagement and adherence to treatment plans.

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