QUALITY OF LIFE EVALUATION IN CELIAC PATIENTS FROM SOUTHERN BRAZIL

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ABSTRACT - Background - Restrictions imposed by the gluten-free diet generate large changes in the daily habits of the celiac patient, causing a negative impact on quality of life. Objective - This study aimed to evaluate the quality of life of patients with celiac disease on a capital in Southern Brazil. Methods - Patients older than 18 years were included, with confirmed celiac disease for at least 60 days in the period from June to October 2013. A validated questionnaire, with specific questions to assess the patient’s quality of life celiac was applied. A total score ranged from 20 to 100 points; the higher the score, worse quality of life. Results - A total of 103 questionnaires were evaluated, 96 (93.2%) female, with average score 56.6±12.35 (28 to 88 points). The comparison between the questionnaire scores and family income was not significant (P=0.139). Patients diagnosed less than 1 year have poorer quality of life than those with more than 10 years (P=0.063). Patients older than 60 years had better quality of life compared with the younger ones (P=0.04). Conclusion - There was no association between quality of life and factors such as family income, length of diet and age at diagnosis. Chronological age greater than 60 years has positively influenced the quality of life of celiac patients.


INTRODUCTION

Celiac disease (CD) is characterized by a chronic inflammatory response in the intestine in response to ingestion of gluten present in the diet. It is a multifactorial disease, which leads to intestinal inflammation, loss of intestinal villi and, therefore, nutrient malabsorption, that results from the combination of genetic susceptibility, environmental exposure and immune response(5). The classic form of the disease presents with gastrointestinal symptoms, such as bloating and chronic diarrhea, but atypical forms are frequent and range from extra-intestinal symptoms until asymptomatic forms(12, 13). CD primarily affects individuals with European descent and its development is strongly influenced by genetics. In Brazil, the South and Southeast regions have the highest prevalence, both for its strong European colonization and the greater availability of diagnostic tests.

An international multicenter study reported that the prevalence of CD worldwide is approximately 1%, with high variation between countries. The prevalence is estimated to be increasing, although the disease is often not diagnosed(11). To estimate the prevalence of CD in general population, blood donors have been investigated. In Brazil, there are currently studies of blood donors in Brasilia, Curitiba and Ribeirão Preto showing a variation in prevalence between 1: 681, 1: 417 and 1: 214, respectively. These findings indicate that CD should not be considered a rare disease in the country(9, 15). Another finding in studies of epidemiology is that CD is most prevalent in women, with an estimated ratio of two women patients for each man(12, 13).

Early diagnosis and treatment are essential in preventing complications such as malnutrition, osteoporosis, infertility and cancer(16). Serological tests with antiendomysium antibody and anti-tissue transglutaminase exhibit high sensitivity and specificity (>90%) and are important in the screening of potential celiac patients(18).

Currently, the only available treatment is a gluten-free diet (GFD)(4, 17), which, once established, stops clinical symptoms, combats nutritional deficiencies, and reduces the risk of developing other autoimmune diseases and malignancies associated with CD(21). The restrictions imposed by the GFD generate large changes in the daily habits of the patient and their families and can be difficult to maintain throughout life(11, 22). Thus, there are studies that show an adequate response to therapy in over 30% of patients(14, 14). The difficulty of finding gluten-free
foods, the high cost and the inadvertent consumption of food containing the substance are factors that contribute to lower adherence to the diet\(^3\).

New treatments for CD have been studied, which include the development of transgenic grain gluten free, blocking epithelial zonulin receptor, and vaccines that induce oral tolerance to gluten. These types of treatment are encouraging results in phase II and III studies and do not depend on patient compliance to the diet\(^3\).

CD, like most chronic diseases, has a negative impact on the quality of life (QOL) of the patient, both for its clinical manifestations that are variable for each individual, and by food restriction that complicates, among other things, interpersonal relationships\(^8\).

The present study aimed to evaluate the QOL of patients with CD on a capital in southern Brazil through validated questionnaire, comparing the results with clinical and epidemiological data.

**METHODS**

The study was observational analytic cross-sectional design and was approved by the Ethics Committee on human research of Positivo University.

Patients older than 18 years were included, with a confirmed diagnosis of CD for at least 60 days, attending meetings of the Association of Celiac from Paraná (ACELPAR) and signed an informed consent form (ICF). Individuals who did not meet these criteria were excluded.

The ACELPAR aims to inform about the CD, providing support, recipes and meetings that are held monthly.

Two types of questionnaire were used in the period from June to October 2013. The first with clinical and epidemiological questions (chronological age, age at diagnosis, gender, ethnicity, nationality and family income) and the second entitled Celiac Disease - Quality of Life (CD-QOL)\(^7\), composed of 20 specific questions to assess the patient’s QOL celiac, validated in English and translated into Portuguese by the authors of this study. Each question was rated from one to five, and the total score ranged from 20 to 100 points; the higher the score, the worse QOL.

**RESULTS**

Over 103 questionnaires were evaluated, 96 (93.2%) patients were female and seven (6.8%) were male. Because of the low number of men who responded to the questionnaire, its exclusion from the statistical analysis was required. Demographical data are shown in Table 1. All respondents reported being doers of GFD.

Among the 96 questionnaires analyzed, the average score was 56.63 ± 12.35 (95% CI = 53.1 to 58.9 points), with a minimum value of 28 and a maximum of 88 points. Twenty-four patients (25%) had higher scores on average and a standard deviation of 68 points; of these, eight (8.3%) totaled more than 80 points, suggesting a worse QOL. Considering that 75% of patients accounted for less than 68 points, it was inferred that the QOL assessed by the questionnaire used in the study was fairly well studied in CD.

The comparison between the CD-QOL scores and family income (Figure 1) lower and higher than five minimum wages was not statistically significant (\(P=0.139\)) and less than five times and greater than ten times the minimum wage (\(P=0.100\)).

On average, the time diagnosis was 5.1 years, ranging between 1 and 34 years. It is noteworthy that the time of diagnosis implies GFD time for most patients. When com-

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**TABLE 1. Demographic data of the patients studied (n=103)**

<table>
<thead>
<tr>
<th>Age</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average 42.1 ± 11.0</td>
<td>16 (16.6%)</td>
</tr>
<tr>
<td>18-30 years</td>
<td>48 (50%)</td>
</tr>
<tr>
<td>31-45 years</td>
<td>21 (21.8%)</td>
</tr>
<tr>
<td>&gt;61 years</td>
<td>11 (11.4%)</td>
</tr>
</tbody>
</table>

**Ethnicity**

<table>
<thead>
<tr>
<th>Ethnicity</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>White</td>
<td>89 (92.7%)</td>
</tr>
<tr>
<td>Afrodescendant</td>
<td>5 (5.2%)</td>
</tr>
<tr>
<td>Yellow</td>
<td>2 (2%)</td>
</tr>
</tbody>
</table>

**Family income**

<table>
<thead>
<tr>
<th>Class</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class A and B</td>
<td>42 (43.7%)</td>
</tr>
<tr>
<td>Class C, D and E</td>
<td>33 (34.4%)</td>
</tr>
<tr>
<td>Not answered</td>
<td>21 (21.9%)</td>
</tr>
</tbody>
</table>

**FIGURE 1.** Scores obtained on Celiac Disease - Quality of Life and family income in celiac disease patients. SM: minimum wage.
paring the CD-QOL score and the time to diagnosis of CD lower and higher than four years (Figure 2), no statistical difference ($P=0.28$) was noted. Patients diagnosed less than 1 year were noted to have poorer QOL than those with more than 10 years of diagnosis, with no statistical significance but pointing trend ($P=0.063$). The comparison between participants who obtained the lowest and highest CD-QOL score at the time of diagnosis also did not reach statistical significance ($P=0.74$).

**DISCUSSION**

QOL evaluate objective and subjective aspects of everyday life. It also evaluate how the needs and expectations of patients are affected by their condition. From this perspective, some authors investigated the impact of CD, and also the GFD, on patients lives\(^4,22\). However, no studies on the topic were found in Brazil.

The proportion of CD between the genders is two women to one man\(^22\). However, in this study, it was necessary to remove the statistical analysis of the data provided by men because of their low adhesion to the research. This fact finds that the low participation of men in ACELPAR meetings, where the questionnaires were administered, and more broadly, their lowest interest in health care, when compared with women. Other authors reported similar findings\(^6,20\).

Importantly, the ethnic difference between Southern Brazil and other regions of the country was due to the strong European colonization present in the area of data collection. CD is a disease classically associated with white ethnicity and highlights the presence of celiac who declared having African (4.8%) and Asian descent (2.9%), showing a strong ethnic mix of Brazil, even in the South.

In 2011, Barrat et al.\(^4\) assessed the likelihood of developing anxiety and depression in patients with CD. These authors showed that the celiac in the UK have a reduced QOL and are more likely to develop anxiety and depression compared with controls of the same age and gender. In disagreement with the results reported, the present study found that most patients did not have depressed mood.

In 2007, Häuser et al.\(^10\) reported that the correlation between the impact of CD and GFD in the QOL of patients...
is conflicting. American, Canadian and Swedish studies reported similar health-related findings between celiac and the general population with QOL; Italian, German, and Irish studies have shown a reduced QOL. Characteristics related to cultural habits and sociodemographic characteristics partly explain these differences between countries. On the other hand, the results of studies on the QOL of CD cannot be fully comparable because of the various instruments used to collect data with different scores and forms of analysis\(^{(10)}\).

In this study, the average score of CD-QOL was 56 points, from 20 to 100 points, according to calculations; 75% of the respondents had values below 68 points, suggesting that the QOL of patients evaluated is slightly better than that found in the literature.

Zarkadas et al.\(^{(22)}\) demonstrated the important role of celiac organizations who offer emotional and social support and encourage adherence to the diet. Häuser et al.\(^{(19)}\) noted the dissatisfaction of patients in relation to the amount of information provided by the physicians and explained the importance of good physician-patient relationship, which also improves adherence. In the present study, the majority of patients were considered to be sufficiently informed about CD. It is possible that the support of ACELPAR and the medical care offered to patients assessed have influenced the results.

The perception that gluten-free products are generally more expensive than the others suggests that household income may interfere with the GFD. Barrat et al.\(^{(4)}\) reported that patients with higher socioeconomic status and college education are more adherent to the GFD because they have more resources to do so. However, there is no evidence to show a clear correlation between these factors and better QOL in patients with CD. In the present study, no statistically significant association between family income and the CD-QOL scores was observed. Additionally, other factors related to the GFD could influence the QOL of these patients. One of the questions this study asked about was the general and constant concern. The dissatisfaction of patients in relation to the amount of information provided by the physicians and explained the importance of good physician-patient relationship, which also improves adherence. In the present study, the majority of patients were considered to be sufficiently informed about CD. It is possible that the support of ACELPAR and the medical care offered to patients assessed have influenced the results.

Assessing the chronological age of the surveyed, better QOL of patients older than 60 years, when compared with other age groups, was found. It is possible that factors related to this phase of life lead that improved QOL among them stands out to the best adaptation to GFD, more meals taken at home and more time available for food preparation. Local features such as greater access to sources of food without gluten and in greater ability and experience in food preparation are a positive influence and improve adaptation to GFD.

Limitations encountered in the study stemmed from the fact that there is not a record of the number of celiac patients in Brazil, not allowing the calculation of the sample. Similar limitation has been reported by Zarkadas et al.\(^{(22)}\) in Canada. In addition, people who frequent celiac associations have greater interest and are better informed about the disease. They also, receive and exchange experiences on how to adapt to the GFD, and these factors can positively influence QOL. Moreover, questions of an intimate nature are not subject to this bias. We also lament the fact of not being able to make comparative analysis between the genders, due to the low number of male respondents to the questionnaire. It is noteworthy that celiac patients should receive a comprehensive treatment that addresses the psychosocial and not just the biological aspects so that the disease causes little influence on their QOL.

The present study showed that the QOL of patients interviewed in a capital of Southern Brazil is higher than expected. There was no association between QOL and factors such as family income, length of diet and age at diagnosis. CD patients surveyed felt well informed, showed no declining trend and showed no constant worry about their food. It is believed that these results are linked to the information and support received, reinforcing the importance of patients associations to better QOL for celiac. Finally, it was found that chronological age greater than 60 years has positively influenced the QOL of celiac patients.

**Acknowledgment**

The authors appreciate the cooperation of all members of ACELPAR (Associação dos celíacos do Paraná; http://www.fenalcebra.com.br/acelpar/institucional/), in particular the president Juliana Gomes Werneck.

**Authors’ contributions**

Castilhos AC, Kotze LMS and Nisihara RM: conception and study design analysis and/or interpretation of data; final approval of manuscript, writing of the manuscript or review of its content. Gonçalves BC Lanzoni LA, Metzger LR and Silva MM: data collection, analysis and interpretation of data; approval of the final version of the manuscript.
Castilhos AC, Gonçalves BC, Macedo e Silva M, Lanzoni LA, Metzger LR, Kotze LMS, Nishihara RM. Quality of life evaluation in celiac patients from Southern Brazil

RESUMO - Contexto - Restrições impostas pela dieta isenta em glúten podem gerar grandes mudanças nos hábitos diários do paciente celiaco, causando um impacto negativo na sua qualidade de vida. Objetivo - Este estudo teve como objetivo avaliar a qualidade de vida de pacientes com doença celiaca, em uma capital do Sul do Brasil. Métodos - Pacientes maiores de 18 anos foram incluídos, com doença celiaca confirmada há mais de 60 dias, no período de junho a outubro de 2013. Um questionário validado, com perguntas específicas para avaliar a qualidade de vida do paciente celiaco foi aplicado. A pontuação total nesse questionário varia entre 20 a 100 pontos; quanto maior a pontuação, pior a qualidade de vida. Resultados - No total 103 questionários foram avaliados, sendo 96 (93,2%) do sexo feminino, com pontuação média de 56,6±12,35 (28-88 pontos). A comparação entre os escores do questionário e renda familiar não foi significativa (P=0,139). Pacientes diagnosticados há menos de 1 ano, apresentam pior qualidade de vida do que aqueles com mais de 10 anos (P=0,063). Pacientes com mais de 60 anos apresentaram melhor qualidade de vida em comparação com os mais jovens (P=0,04). Conclusão - Não houve associação entre a qualidade de vida e fatores como renda familiar, tempo de dieta e idade no momento do diagnóstico. A idade cronológica superior a 60 anos influenciou positivamente a qualidade de vida de pacientes celiacos.

DESCRITORES - Doença celiaca. Qualidade de vida. Dieta livre de glúten.

REFERENCES
