Background

In recent decades, the prevalence of diabetes has risen sharply around the world. According to estimates of the International Diabetes Federation, there are currently 451 million people with diabetes aged 18–99 in the world, and it is predicted that by 2045, this number will increase to 629 million [1]. Is this increasing number of people with diabetes getting more and more understanding and social support? Can any person suffering from diabetes always and everywhere admit to being diabetic?

In the literature, we encounter the problem of stigmatization and discrimination of people with diabetes. People who do not suffer from diabetes assume that diabetes is not a stigma [2]. In contrast, people with diabetes report that stigmatization is a serious problem for them – one which they experience in different areas of life, especially in the workplace or in relationships [2].

In the U.S., a study was carried out to measure the stigma of diabetes and the associated psychosocial effects within a large population of patients, using an online survey sent to 12,000 people with diabetes [3]. The results reveal that the majority of respondents with type 1 diabetes (76%) or type 2 diabetes (52%) reported that diabetes is associated with stigma [3].

The Second Diabetes Attitudes, Wishes and Needs (DAWN2) study, which covered over 8,500 adult diabetics from 17 countries, showed that 17.6% of all patients thought their affliction generated discrimination, intolerance and a lack of support from the communities in which they lived [4]. In Poland, the percentage of patients declaring similar experiences was higher and amounted to as much as 24% [4]. It is disturbing that even the very fact of becoming diabetic may cause shame in some people [5]. Such feelings of shame can prevent the afflicted from informing people within their social environment about the illness. This can have a negative impact on the course of diabetes therapy.

In the literature devoted to the problem of informing about the disease, we find, first of all, issues concerning the manner in which medical personnel inform the afflicted about the course of their disease. The intent of the provided information is to compel the patient to adopt their doctor’s recommendations, and, consequently, to ensure effectiveness of the therapy. It is also emphasized that information of this nature, regardless of what disease it concerns, creates a specific doctor–patient relationship and builds the patient’s trust [6] and their willingness to care for their own health.

The problem of providing crucial information to the patient’s family, is considerably less often taken [7]. When done correctly, doing so fulfills similar goals, because the patient expects understanding, support and acceptance from their family members. Indeed, the importance of the family has an additional dimension as the family has significance with regard to medical care and health protection, especially in chronic diseases, as it partners the health care system in performing traditional health-related tasks [8].

In everyday diabetological practice, however, we encounter situations wherein patients signal that they do not always admit to having diabetes. It is likely that in such scenarios, the
afflicted fear potential discrimination or stigmatization should they admit to being diabetic. Analyses of the information that the ill person communicates with their social environment are only occasionally performed/carried out. The only material available in the literature are individual studies on the concealment or disclosure of information about the disease by patients with diabetes, especially in the workplace [9–11]. Thus, there is a need to understand the social implications of being diabetic.

Objectives

The aim of the study is to understand why those afflicted with type 2 diabetes inform only certain individuals within their social circles, of the course of their disease. An attempt was also made to estimate the relationship between the fact of limiting admission of being diabetic and the values of the selected parameters of metabolic diabetes control.

Material and methods

Setting and participants

The study covered 136 patients with type 2 diabetes, including 71 women and 65 men (age – median 62.5, min–max 40–84) who reported to the Diabetological Outpatient Clinic of the Chair and Department of Family Medicine, Medical University of Lublin, for a medical appointment within a 6 month period, i.e. from September 2007 to February 2008. The study used a previously developed original questionnaire assessing the socio-economic and family situation of patients with type 2 diabetes [12]. The respondents were asked to indicate people from their social environment (family members, friends, neighbours and others) who they inform about their illness. The respondents also gave reasons for telling others about their disease. The categories of causes included the expectation of greater emotional support, maintenance of dignity, as well as the health, social and professional safety of the patient. The metabolic control of diabetes was assessed based on the level of A1c glycated haemoglobin (HbA1c) and fasting plasma glucose in the venous blood. Satisfactory values of fasting glucose was assumed to be ≤ 110 mg/dl and for HbA1c ≤ 7%.

The study design received a positive opinion from the Bioethics Committee (KE-0254/116/2007).

Statistical methods

The statistical analysis was performed using the IBM SPSS program version 25. The Pearson χ² test was used in order to determine the statistical significance of the correlation between the variables. Values below 0.05 were assumed to be significant. The Kolmogorov–Smirnov test with the Lilliefors correction was used to ascertain normal distribution, herein, the confidence in the Kolmogorov–Smirnov test with the Lilliefors correction was set at the average of 95%. Age did not have normal distribution, but variables such as duration of diabetes, fasting interval was set at the average of 95%. Age did not have normal distribution, but variables such as duration of diabetes, fasting interval was set at the average of 95%. Age did not have normal distribution, but variables such as duration of diabetes, fasting

Results

Participants and descriptive data

Persons aged < 65 years accounted for 55.9% (76) of the respondents, and those aged ≥ 65 years – 44.1% (60). The vast majority of the respondents lived in the city (75.7%; 103) and the remaining 24.3% (33) in the countryside. The percentage of the respondents living with at least one person in a household was 82.4% (112), and those living alone – 17.6% (24). Over 2/3 of the respondents (67.4%; 91) were married and the remaining 32.6% (44) were unmarried. The group characteristics are presented in Table 2.

People with vocational and secondary education dominated among the respondents (66.9%; 91), 18.4% (25) had primary and 14% (19) – higher education. Most of the respondents were professionally inactive (72.1%; 98). The people from the professionally active group were divided into employed (84.2%; 32) and self-employed (15.8%; 6). The mean duration of diabetes in the study group was 8.72 years (SD = 6.898).

The mean fasting blood glucose level in the study group was 122.22 mg/dl (SD = 31.471). In 61% (83) of the subjects, the fasting blood glucose level exceeded 110 mg/dl. The mean HbA1c level found in the study group was 6.6% (SD = 6.599). Satisfactory HbA1c values (≤ 7%) were reported in 78.7% (107) of the subjects.

Main results

It appears that there are four groups of people who are most often informed by patients with type 2 diabetes with regard to their illness: 1) family members – both those who live with the patient (99.1%; 111) and those who do not live with the patient (86.0%; 117), 2) other people with diabetes (80.1%; 109), 3) friends (72.8%; 99) and 4) neighbours (63.2%; 86) (Figure 1). People at workplace are less frequently informed about the disease. Every second employed respondent (50%; 16) provided his/her employer with the information about the disease; a slightly larger percentage of patients (57.3%; 18) inform their workmates. With regard to their disease, all persons living alone in a household with at least one person rarely inform family members not living with them about their illness (100% (24) and 83% (93), respectively, p = 0.03). People who are professionally inactive more often informed their friends about the disease than did those professionally active (77.6% (76) and 60.5% (23), respectively, p = 0.045).

Neighbours are an important category of people informed by patients about their illness. The variables which differentiate the frequency of informing neighbours include: 1) sex – more often women than men (71.8% (51) and 53.8% (35), respectively, p = 0.03), 2) age – more often older people aged ≥ 65 years than those younger (75% (45) and 53.9% (41), respectively, p = 0.011),
than women expect understanding of their situation (56.9% (37) and 39.4% (28), respectively, \( p = 0.041 \)). Similarly, older patients (≥ 65 years) more frequently than younger mentioned the willingness to be understood by people they inform about the disease (60% (36) and 38.2% (29), respectively, \( p = 0.011 \)). As for the needs related to diabetes therapy, the situation is similar. the older the person, the more willingly he/she speaks about his/her illness so that others would not hinder their compliance with medical recommendations (60% (36) and 38.2% (29), respectively, \( p = 0.011 \)). Also more often, older patients who inform others about their disease expect help in compliance with medical recommendations (46.7% (28) and 19.7% (15), respectively, \( p = 0.001 \)). People who are professionally inactive more frequently than those professionally active inform others about their health problem in the hope of receiving help in complying with medical recommendations (37.8% (37) and 15.8% (6), respectively, \( p = 0.013 \)). People living in a household with at least one person, as opposed to those living alone, expected to be relieved of some duties due to their affliction (15.2% (17) and 0%, respectively, \( p = 0.041 \)).

no statistically significant relationship was demonstrated between the fact of informing about the disease and education, duration of diabetes and fasting glucose level.

### Discussion

**Key results and interpretation**

The results of the study confirm literature data that the family is the most important source of support for diabetic patients.
Lawksa et al. also showed that the majority of diabetics expect help from their family [16]. The near family is considered the group of the greatest trust with regard to freely informing about the affliction without the expectation of negative reactions. Indeed, Stopford et al. concluded that the family support is most often associated with the reduction of HbA1c level [14].

The study conducted by Mohebi et al. also revealed that patients with higher HbA1c values felt less social support [17].

Another large group (80%) informed about their disease are other people with diabetes. The analysis of reasons for informing such about diabetes has shown that more than half of patients shared the information about their illness with other patients based on the rule of reciprocity. The exchange of experience and mutual assistance between patients with diabetes may, on the one hand, improve patients’ self-esteem, and on the other hand, expand their knowledge about treatment, self-care and coping strategies.

It was also demonstrated that friends and neighbours are important social groups. Almost 2/3 of all patients with diabetes trust their neighbours and tell them about their illness. These are more often women, older people aged ≥ 65 years old, unmarried persons and those professionally inactive. It seems that with time, neighbourhood ties tighten and neighbours can be an important source of support for patients in the treatment of type 2 diabetes. On the other hand, unmarried persons more often than married inform neighbours about their illness hoping for support. Shaw et al. demonstrated that neighbours and neighbourhood resources seem to have a great impact “on adherence to diabetes self-care behaviours” [13].

The study found that the main reason why patients with type 2 diabetes informed others about their illness was a desire to prove that “you can live a normal life with diabetes”. This belief is more common among patients with well-controlled diabetes (HbA1c ≤ 7%) who set a good example, and, in this way, can convince other patients that diabetes can be overcome and that they enjoy all areas of life. It was also demonstrated that, on average, every second patient with diabetes informs others about their illness, expecting help in case of hypoglycaemia, understanding of the disease and non-hindrance in their compliance with medical recommendations or is motivated by a willingness to share experience and knowledge about diabetes.

Interestingly, patients with poor glycemic control (HbA1c > 7%) far more often than patients with good glycemic control (HbA1c ≤ 7%) inform their social environment about their illness, expecting help in case of hypoglycaemia. This may be an important hint for doctors dealing with the therapy of diabetic patients that high HbA1c levels in these patients may be due to fear of hypoglycaemia and to a deliberate maintenance of glycemia at a higher level. This is especially true for workplaces where the fainting of a sick employee with diabetes (e.g. drivers) may endanger other people in a different way. Ruston et al. also showed that diabetic patients were reluctant to disclose their illness at work and reported the need for support if they were stigmatized or treated inappropriately, e.g. if symptoms of hypoglycaemia occur. He stated that diabetic employees strive to maintain glycemic values at a higher than recommended level to prevent hypoglycaemia or exposing oneself to the development of chronic diabetes complications [22].

Limitations of the study

The results of our study concern a group of patients with type 2 diabetes, characteristic of the older age category, in a large percentage of the professionally passive. This does not give a full orientation in the problems of informing about the disease by a larger group of patients with diabetes with more diversified demographic characteristics and professional and family situations. Future research should be extended to patients with type 1 diabetes, and in younger age categories, including a larger group of professionally active patients.

Conclusions

1. Patients with type 2 diabetes most often inform their family and people with diabetes of their situation, and the least frequently the employer.
2. The main reason for informing others about the illness is a desire to prove that “you can live a normal life with diabetes”.
3. There is a negative correlation between the HbA1c level and a willingness to prove that “you can live a normal life with diabetes”.
4. There is a positive correlation between HbA1c, and the expectation of receiving help in case of hypoglycaemia.
5. Family physicians should encourage patients with diabetes to inform others about their illness, especially in workplaces, both for their own safety and for the safety of people in their environment.
6. Failure to inform the social environment about the diagnosis of diabetes can have negative consequences for the health and life of both the patient and people from the social environment. The reasons for this behavior require further research.

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References


