HEALTH BEHAVIOUR CHANGES OF CUTANEOUS MELANOMA SURVIVORS IN SLOVENIA - A QUALITATIVE STUDY

SPREMEMBE V ZDRAVSTVENEM VEDENJU BOLNIKOV Z MELANOMOM KOŽE V SLOVENIJI - KVALITATIVNA RAZISKAVA

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ABSTRACT

Introduction: Most data related to cutaneous melanoma survivors’ health behaviour comes from epidemiological studies and is predominantly concerned with safe-sun behaviour and self-examination. Data regarding other changes of health behaviour are scarce and so are qualitative studies in this realm. The aim of our research is to acquire insight into the experiences of patients with cutaneous melanoma in Slovenia. How did they react to the diagnosis, which changes did they introduce in their health behaviour and how do they assess the role of family doctors?

Methods: Using the qualitative approach of collective case reports, a demographically diverse group of patients with different forms and stages of cutaneous melanoma was selected. Semi-structured interviews conducted by a psychologist were recorded and transcribed verbatim. For data processing, the approach of Qualitative Content Analysis was applied.

Results: We integrated interviewees’ experiences after the diagnosis of cutaneous melanoma in several subcategories: either they did not introduce any changes or they mentioned changing their habits when exposed to the sun and performing skin self-examination; they also emphasized their ways of dealing with stress and raising awareness about melanoma among family members and friends. The role of family doctors in the prevention and care appears unclear; even contradictory.

Conclusions: We obtained insight into the experiences of Slovenian patients with cutaneous melanoma. The interviewees prioritised safe behaviour in the sun, strengthening of psychological stability and raising awareness about melanoma. Findings will be used in the creation of a structured questionnaire for national epidemiological survey.

IZVLEČEK

Ključne besede: melanom kože, obvladovanje stresa, izkušnje, združenost, kvalitativna raziskava

Uvod: Večina podatkov, povezanih z zdravstvenim vedenjem bolnikov z melanomom kože, je pridobljena z epidemiološkimi raziskavami. Raziskave se pretežno ukvarjajo z varnim obnašanjem na soncu in samo-pregledovanjem kože. Bistveno manj poznamo druge spremembe v zdravstvenem vedenju teh bolnikov, prav tako je – tudi v tujih literaturi – le malo kvalitativnih raziskav na to temo. Namen raziskave je pridobiti vpogled v izkušnje bolnikov v Sloveniji po diagnozi melanoma kože; kako so se odzvali na bolezen, kakšne spremembe so vpeljali v svoje (zdravstveno) vedenje in kako vidijo vlogo osebnih zdravnikov.

Metode: Ustrezno raziskavo je pripravljalo skupino bolnikov različnih spolov, starosti, izobrazbe in telesnih oblik izbereti. Znesek bolnikov je bil prikazan kot različne podkategorije: bodisi intervjuvanci niso namenili sprememb v zdravstvenem vedenju, bodisi so prevedli na neposredno podporo v zdravstvenem vedenju, bodisi so prihajali v več podkategorije: bodisi intervjuvanci so izvedli sprememb ali pa so obstajali spremembe pri začetki pred soncu in samopregledovanju kože; pojavljuje se tudi odziv na stres in osveščanje o sebi glede melanoma. Sredine, kjer se prihajalo javnosti bolnikov, so medtem postale boljši ali pa spremenili svoje vzorce zdravstvenega vedenja. Bolniki so se izpostavili bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost, več bolnikov se je izpostavilo bolniki v telesno aktivnost.

Zaključek: Pridobljen je pričakovani vpogled v izkušnje slovenskih bolnikov z melanomom kože. Intervjuvanci so izpostavili varno obnašanje na soncu, krepitve čustvene stabilnosti in ozaveščanje bližnjih glede melanoma. Izklopka bodo uporabljeni pri oblikovanju strukturiranega vprašalnika za epidemiološko raziskavo, ki bo ovrednotila spremembe v zdravstvenem vedenju bolnikov z melanomom kože in iskala dejavnike, ki ključno vplivajo na telesne aktivnosti. Velik del intervjuvancev je ljudi okoli sebe pričel osveščati glede melanoma kože, tveganih bolnikov v Sloveniji po diagnozi melanoma kože; kako so se odzvali na bolezen, kakšne spremembe so vpeljali v svoje (zdravstveno) vedenje in kako vidijo vlogo osebnih zdravnikov.
INTRODUCTION
Cutaneous melanoma is one of the most common malignancies in the world. Given the possibility of aggressive metastases, it is the most dangerous form of skin cancer (1). Both the incidence (recorded or estimated) and the mortality of cutaneous melanoma are significantly higher in the more developed parts of the world (2).

In IARC estimations for 2018, are among 20 countries most heavily burdened with melanoma, as many as 16 European countries; Slovenia (with estimated ASR 18.6) occupies the 10th place in the world and 8th in Europe (3). In the period 2011 to 2015 an average 124 persons per year died because of cutaneous melanoma in Slovenia, with the average age-standardized mortality rate in this period being 2.85 (3.50 for men and 2.33 for women) (4). 5-year survival rate in Slovenia, which was at the beginning of the 21st century 80.4% and worse than European average (5), has substantially improved to 89.4% in the 2011-2015 period. In the last twenty years much has been said about melanoma, but usually among professionals or in the direction from profession to population. Nowadays, these processes of communication and awareness are well established, either through everyday medical procedures or through prevention campaigns (6). However, when speaking with patients in dermatology clinics in Slovenia, we still notice much fear and little actual knowledge about melanoma (personal communication, 7).

The stage at which melanoma is detected and surgically removed is the most important factor for the prognosis of the disease and survival; prevention on primary and secondary level remains the key factor for the reduction of melanoma-rated mortality (8-12). This is especially important for melanoma survivors, who have a significantly increased (13) risk of acquiring second primary melanoma and should be repetitively counselled about the risk and about the importance of preventive health behaviour. To optimize our messages and interventions regarding preventive behaviour, it is crucial to (i) recognize what has changed in health behaviour after the diagnosis of melanoma and (ii) to understand the triggers to the changes. In the literature there is relatively little data (9-11, 14-16) regarding the health behaviour of melanoma survivors. For the Slovenian population we have none. The aim of this qualitative part of our research is to understand how melanoma survivors in Slovenia percepts the disease, how do they cope with it and how do they assess the role of the family doctor. Its key objective is the identification of changes in health behaviour after obtaining the diagnosis: what they do, what matters to them and whether they mention anything we have not thought about or learned from the literature.

Collected data is going to be used in subsequent epidemiological study (17), results of which shall help in the planning of future public health interventions in Slovenia.

METHODS
We performed a qualitative study of patients’ experiences before and after the diagnosis of cutaneous melanoma; we used the approach of the collective case report. Among different approaches in qualitative research methodology, case studies are recommended when complex phenomena are to be studied within their contexts (18-20). They are the most powerful when answers on «how» and «why» are sought. Based on the literature, the collective (multiple) case studies enable the researcher to detect and explore the differences through replication of findings across cases (19-21). Regarding this, cases should be selected primarily because of their own intrinsic value (22). However, to better understand the underlying phenomenon, parallel sampling design is often applied: «the selected cases are treated as a set and their voice is compared to all other cases one at a time in order to understand better the underlying phenomenon, assuming that collective voices generated by the set of cases lead to data saturation» (23).

2.1 Participants
Altogether, ten patients from a dermatological outpatient clinic in Ljubljana, Slovenia were included (5 men, 5 women). We invited them to participate in the study during their regular medical check-up as melanoma survivors, however, they came to the interviews on the agreed term in their free time. Inclusion criteria were as follows: (i) diagnosis of cutaneous melanoma in the past (single, multiple, invasive or in situ) (ii) age >18 years and (iii) fluent in Slovenian language. To ensure the anticipated group diversity of parallel purposeful sampling, we invited patients from different geographical regions, educational background, age, gender and different melanoma stages. Parallels were gender and whether an interviewee had had one or multiple melanomas. In terms of time from the diagnosis, patients were limited only to the fact that at least one year had passed since the (first) melanoma. Group characteristics are depicted in Table 1. Patients were gathered in period from January 2015 until April 2015.
To ensure reliability of the coding, two independent researchers coded the texts. When applied to their coding, the coefficient of agreement was high (97%). Regarding the validity, when a coding frame is mostly “data driven”, as it happens in this research, “face” validity is advised (21). Validity is a measure of assessment between concept of the study and its procedure, and face validity displays the “extent to which your instrument gives the impression of measuring what it is supposed to measure” (21). The assessed face validity of the data was high, as there were no abstract or residual categories left.

According to the leading questions of the interview, data were assessed as important/relevant and classified into relevant categories.

3 RESULTS

Data based on interviews with 10 patients were categorized according to leading questions. As our aim was identification of experiences of patients who have been diagnosed with cutaneous melanoma, in the foreground were their reactions on the disease and changes in health behaviour they implemented.

Exploring the responses of the interviewees in terms of changes in their health behaviour led to the main category “impact of being diagnosed with cutaneous melanoma on health behaviour”.

Patients’ answers are depicted in Table 2.

### Table 1. Characteristics of the participants.

|                          | Total (men/women) |
|--------------------------|-------------------|
| Participants (n)         | 10 (5/5)          |
| Age, mean (years)        | 51.8 (62/41.6)    |
| Diagnosis *              |                   |
| MIS                      | 8 (6/2)           |
| SSM                      | 19 (10/9)         |
| NM                       | 1 (1/0)           |
| LM                       | 1 (1/0)           |
| One MM                   | 4 (2/2)           |
| Two or more MM/MIS       | 6 (3/3)           |
| Duration from (last) diagnosis to interview, mean (years) | 3.7 (2.6/4.8) |

*melanoma in situ (MIS), superficial spreading melanoma (SMM), nodular melanoma (NM), lentigo maligna melanoma (LM)*
Table 2. The impact of being diagnosed with cutaneous melanoma on health behaviour.

| Main categories                | Main subcategories | Detailed subcategories                                      | Codes                      |
|--------------------------------|--------------------|-------------------------------------------------------------|----------------------------|
| Health behaviour management    | No changes         | living the same way                                         |                            |
|                                | Changes in sun behaviour | use of protective creams                                | -different- creams          |
|                                |                     | higher SPF used                                            | physical protection        |
|                                |                     | vigilance in hours                                          | avoiding the sun           |
|                                | Changes in skin self-examination | checks regularly                                  | checks occasionally         |
|                                |                     | checks before the medical check-up                        | being more attentive to the skin |
|                                | Dealing with the stress | strengthening of mental stability                        | independently              |
|                                |                      | with help of an expert                                     | conversation with family/friends |
|                                |                      | physical activity                                          | stroll                     |
|                                | Raising awareness of melanoma | among kinship                                         | among acquaintances         |

Several subcategories emerged. Two interviewees reported that they have not changed anything important: “I do not think that I did [change anything]. Maybe I have sorted certain things out, but basically I live exactly the same, because I was already convinced that I did not make any mistakes” (male, 55).

The rest of the subcategories were about changes in different aspects of health behaviour, where the first two, focusing on participants’ attitude regarding the UV exposure and their examination of the skin and eventual changes, were expected, and they were distinctly realized in the interviews. The third subcategory is mainly speaking about coping with the stress of malignant disease: trying to (re)gain inner peace, either with work on one’s mind/soul or with increased physical activity to alleviate the stress.

The last subcategory is dedicated to the care for (the important) others, raising awareness about melanoma, safe behaviour in the sun and being observant regarding the changes on one’s own skin.

As expected, a large part of the interviewees became much more cautious when exposed to the sun. Most of them started practising cautious protection from UV rays, either with SPF products or with clothes: “I also wear a swimming cap, I am strictly protected… the most important is thorough protection from the sun, covered (with clothes), cream applied, this is the most important for sure, so to speak the only one…” (male, 74) or they became vigilant regarding the time of the day, when spending time outside “the most important instruction being that one is not exposed to the sun when it is at its strongest…” (male, 55). Some participants even began to avoid being outside during the day: “the fears about the sun exposure… came subconsciously... now I really go to extremes, I meet with friends at 18.00 only…” (female 29).

They also quite often mentioned the use of protective creams with chemical protection factors; either they started using them or they advanced to a higher SPF; one of interviewees started to use an alternative, “different” cream: “…I actually started using creams without any chemistry; for the face and body… there is no chemistry, everything is on mineral foundation; before (the melanoma) there was nothing of this, I thought it was not important…” (female, 47).

The issue of skin (self) examination (SSE) is somehow controversial. Namely, only three of ten participants mentioned SSE spontaneously but, after being specifically asked, all of them asserted that they perform it either
regularly: “I do the examination myself once a month” (female, 47), occasionally, or they are generally more observant regarding their skin: “...since then [melanoma diagnosis], I have been more attentive to the moles” (female, 33).

Outstanding is the majority, who mentioned the need to alleviate the stress connected to diagnosis of cutaneous melanoma. The subcategory of strengthening of mental/psychical stability was addressed most frequently of all categories. To most of the participants sharing their feelings and worries with relatives and friends was helpful: “I talked to my sister... she took a lot of care. Talking to someone who is close to you - at that time there is something gathering inside you - you have to deliver it out of yourself, to talk, to know that you are not alone. That you have somebody you trust” (female, 47). Much more rarely they used different mental techniques or obtained the help of an expert; “...there was so much of everything, I could not sleep...; I started to visit a psychiatrist... I had a feeling that I was insane and I needed a psychiatrist...” (female, 33). Quite a lot of interviewees sought relief in physical activities like yoga, intense strolling and hiking or more strenuous sports: “I am more physically active; I began working out” (female, 47).

The subcategory of raising awareness about melanoma and prevention thereof is represented quite largely - to the same extent as the subcategory about cautiousness towards UV exposure. Most of the participants were focused primarily on family members, while others were even more outspoken: “...in particular I told everybody I knew... in our family the diagnoses were somehow secret until then... so they also started to watch themselves and everybody around me visited a dermatologist” (female, 47).

In the part of the research where interviewees spoke about the role of their family doctors, we designed two main categories: before and after the diagnosis of cutaneous melanoma. All the subcategories were data driven (Table 3).

A weak half of the interviewees mentioned a role of family doctor before the diagnosis of cutaneous melanoma and after it. They assessed the doctor’s attitude before the diagnosis quite critically; sometimes almost angrily. They felt that she/he was not attentive enough to their skin - nobody mentioned that a check-up of the skin lesions was performed and only one participant, with positive family history of melanoma, mentioned that she was advised to self-examine her skin: “...by nature I have many moles and my doctor advised me to check them” (female, 29). However, another participant, phototype 1 and with positive family history of melanoma, felt that total forsaking of skin examination was not appropriate: “...for example, I regularly attend medical check-ups, do the electrocardiogram... but nobody ever mentioned skin self-examination to me” (male, 55). Two participants expressed concern and sounded apprehensive because their family doctor’s diagnostic assessment was wrong, even though they themselves alerted her/him about the lesion: “... a strange lesion appeared... I showed it a few times to my doctor, but she always said: ‘oh, it is nothing important’” (female, 58).

Statements regarding the role of family doctors after the diagnosis of cutaneous melanoma strike as being much less passionate; they sound more as a depiction of a relationship that a patient has with her/his family doctor. Interviewees experienced their family physician’s attitude in diverse ways; as active “...he sent me to blood tests, I am under control...” (female, 47) or passive: “...I believe he is warning the patients... but ... I overtook him with the results...” (male, 74), some experienced it as relieving “...my family doctor also told me that the first level is not that dangerous...” (female, 29) and others as burdensome: “The event... which could shake someone a little... she (the doctor) automatically said, if I were visiting oncology department, if I had psychological support... I saw a question mark above her - why I was not going” (male, 55).

4 DISCUSSION

After they received the diagnosis of cutaneous melanoma, the vast majority of our interviewees introduced changes to their health behaviour. In accordance with before-mentioned data from literature (9-11, 15, 16), they were predominantly changing their habits regarding the sun protection. Only a part of interviewees spontaneously mentioned self-examination although, after being asked directly about the subject, everybody claimed that after the diagnosis they watch their skin more carefully. Reasons
that self-examination was not frequently mentioned are not known. We assume that it is still a common occurrence that people connect prevention against skin cancer only with safe sun behaviour. One of the reasons may be that experts emphasize safe sun behaviour over self-examination in the media and in personal communication with patients (24, 25). However, in Slovenia we heed the guidelines for management of patients with melanoma (26). Patients are rigorously monitored and counselling about safe sun behaviour and skin self-examination are regular parts of medical visits. On the other hand, it may be that patients do not trust in their own capability of recognising an atypical lesion and, as Ajzen stated (27), the limiting factor is low perceived behaviour control. This may well be connected with high fear appeal regarding melanoma in the last few decades and not (yet) strong enough efficacy messages (28).

Interviewees very often mentioned the strengthening of their psychical stability (29); usually this was achieved independently of any professional help (most often through conversation with relatives and friends), only rarely with some help from an expert. Even though there is no lack of evidence that melanoma survivors are in need of psychological support (30, 31), we did not find any data about specific approaches to melanoma patients. Researchers who looked into the cancer treatment in the USA in general, found that, despite the fact that the provision of psychosocial support to the cancer patients was identified as part of good medical care, health professionals often do not detect patients’ need for it. As mentioned in the literature (32, 33), patients are rarely referred to appropriate specialists or treated according to their psychological needs.

Relatively often, interviewees mentioned the intensification of physical activity after being diagnosed with cutaneous melanoma - either outdoors or in the gym. Although the benefits of regular physical activity for melanoma patients have not been investigated, regarding the results of studies, concerning patients with other cancers (34), the benefits are expected and are emphasized in Cancer Fact Sheets (35). Regular outdoor activity after being diagnosed with melanoma is encouraged but, as Lawler and co-workers suggested (36), physical activity and sun protection should be promoted simultaneously.

More than half of our interviewees are raising awareness on the issues of cutaneous melanoma among their relatives and friends. The result is not in concordance with the report from Oliveria and co-workers (16), where even patients who understood the increased risk for family members rarely raised the issue among them.

Surprisingly, considering reports in the literature (37), not even one interviewee mentioned changes in other aspects of healthy lifestyle, like healthy eating or reducing alcohol intake and smoking.

Patients’ experience on the role of family doctors is, strictly speaking, not a component of health behaviour, but already at the first interview the message had such power so that we did not label it as irrelevant; when opinions multiplied, we formed a separate part of the research. Interviewees mentioned the impact of a family doctor in the time before the diagnosis of cutaneous melanoma and, later, after the diagnosis and treatment were performed.

The results are surprising. Almost half of the interviewees think that they belong to a group with a high risk for skin cancer because of their skin type or/and abundance of moles, and two of them are burdened with a family history of melanoma. However, physicians payed no special attention to their skin; the only exception was a warning about the need of skin self-examination to one participant. Moreover, when alerted about a “suspicious” pigmentary lesion (which was later confirmed to be melanoma), the family doctor did not take it seriously.

After the diagnosis of cutaneous melanoma, the situation is somehow fragmented and unconvincing. Half of the interviewees do not even mention a family doctor or her/his response. Others depict different situations; they are experiencing their physician as active, with relieving impact, but also as ignorant and burdening or very passive. Results are consistent with the reporting of Loescher and co-workers (38) who find that communication with people at high risk is inadequate, inconsistent or even inaccurate. Reasons are attributed to (i) the lack of time that a doctor spends with the patient, which is further reduced by all the more extensive administrative requirements, (ii) intensive sub-specialization of doctors and, consequently, lack of knowledge in areas not domiciled and (iii) lack of communication skills. However, as Hajdarević and co-workers (39) stated, healthcare providers should expand the understanding of patients’ experiences and the patterns of seeking medical care. By facilitating access to medical care, early diagnostics will also be facilitated.

4.1 Study Limitations

We are aware of two limitations; interviews were conducted in familiar medical surroundings and half of the interviewees were regular patients of the principal investigator. Those circumstances could have influenced patients’ responses. In order to diminish that influence, a psychologist who is not a member of the staff and was unknown to interviewees conducted the interviews.
Probably we should also mention the number of participants among limitations. We are aware that many qualitative multiple case studies include 30 or more participants but, on the other side, Creswell (19) advises to use four to five cases. Only if there is a rationale does he advise to expand the group. Our rationale was diversity of interviewees. Thus, in the beginning we included 10 patients, with the intention to include more if we do not achieve saturation. However, saturation was achieved by the seventh patient and we saw no reason to expand the group.

Regarding the time interval from the last melanoma, which is different for men and women, we believe that we cannot consider it as a limitation. All interviewees could be labelled as “mature” survivors, and most probably their responses are not an acute (over)reaction to the diagnosis of melanoma but stable changes in health behaviour.

The strength of our study is its design based on the patients’ experience after being diagnosed with cutaneous melanoma and possibility of detailed research of patients’ perception and reaction. Moreover, the research framework was not rigid and when new information emerged the framework could be revised.

Implications for practice: findings served as a background for ongoing quantitative epidemiological survey, which is evaluating the extent of changes in health behaviour and factors that trigger those changes. Collected data will facilitate communication with melanoma patients and help to design corresponding public health interventions.

5 CONCLUSION

Presented qualitative research offers an insight into the experience of patients with cutaneous melanoma in Slovenia and into the changes of their health behaviour. The interviewees set in the forefront safe behaviour in the sun, dealing with stress and raising awareness about melanoma among fellow men. They mentioned skin self-examination less often. The specific role of family doctors in the prevention and care of people with high risk for melanoma appears contradictory or is at least not sufficiently defined.

CONFLICTS OF INTERESTS

The authors declare that no conflicts of interest exist.

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ETHICAL APPROVAL

This study conforms to the recognised standards of the Declaration of Helsinki and was approved by the Republic of Slovenia National Medical Ethics Committee on 13th of May 2014 (reference number 139/05/14).

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