Supportive care needs of patients with rare chronic diseases: multi-method, cross-sectional study

Miram K. Depping1*, Natalie Uhlenbusch1, Yskert von Kodolitsch2, Hans F. E. Klose3, Victor-Felix Mautner4 and Bernd Löwe1

Abstract

Background: In the absence of a cure for the majority of rare diseases, the disease management aims to provide optimal supportive care. The goal of this study was to assess supportive care needs in patients with chronic rare diseases.

Methods: Cross-sectional mixed-method study was conducted using validated self-report scales and open-ended questions to assess supportive care needs. Participants affected by rare diseases across Germany were contacted via patient organizations and centers for rare diseases. N = 304 participants with 81 different rare diseases completed the study, 81.6% were female, mean age was 44.2 years (SD = 12.8, range 16–74). The quantitative results regarding supportive care needs were compared to a reference population of patients affected by cancer (N = 888). Main outcomes were unmet supportive care needs of patients with rare diseases, as assessed by the Supportive Care Needs Survey (SNCS-SF34) and an open-ended question on support wishes.

Results: Patients with rare diseases did not feel sufficiently supported with regard to psychological support, health system and information, physical and daily living, patient care and support, and sexuality needs. The unmet supportive care needs were significantly higher in the patient sample with rare diseases compared to the SCNS-SF34 reference sample of patients with cancer. 60% of patients with rare diseases did not feel sufficiently socially supported.

Conclusions: Patients affected by rare diseases have high unmet support needs in all areas studied. Multidisciplinary care, including psychological support and the provision of information regarding the healthcare system, treatment options, disease course and sexuality, might help address these needs.

Keywords: Supportive care needs, Psychosocial support, Rare diseases, Patient preferences, Patient reported outcomes

Background

300 million people are living with a rare disease worldwide, while every single condition is defined by a prevalence of 5:10,000 [1]. This large group of patients is affected by different somatic conditions, which impact everyday life of patients in various ways. According to a European-wide survey, 8 in 10 patients and informal carers have difficulties completing daily tasks like household chores [2]. Compared to the general population, people living with a rare disease and carers report three times more often to be unhappy and depressed. 7 in 10 patients and carers of patients affected by a rare condition reduced or stopped professional activity due to their family member’s rare disease. These survey data show...
that patients with rare disease face high burden and that healthcare to date does not adequately address the needs of patients with rare diseases, yet. This survey had been carried out by EURORDIS—Rare Diseases Europe (European Organization for Rare Diseases, a non-governmental patient-driven alliance of patient organizations) in want for research of the lives of patients with rare diseases. A review that synthesizes qualitative studies on specific rare conditions, concludes that patients with different rare diseases all face universal burden, such as lack of information [3]. No quantitative studies to date have used validated instruments to assess unmet supportive care needs in different domains across a population of people living with different rare diseases.

Given that there is no cure for the majority of rare diseases, the management of rare diseases aims at providing optimal supportive care. Experiencing burden does not necessitate a want for help, therefore directly measuring patients’ own perceptions of their need for help may be most adequate [4]. The current study sets out to understand what patients with rare diseases wish for above the currently available forms of care. It is the first study aiming at investigating multi-domain supportive care needs in patients with different rare chronic diseases; that is whether they currently feel sufficiently supported with respect to health system and information, physical needs, patient care, sexuality and psychological needs. Supportive care needs in all 5 domains are relevant to a patient’s psychosocial well-being [4]. In order to be able to contextualize the results, we included a comparative sample of patients with other chronic conditions in the analyses, namely patients with cancer. We further explored whether this patient population feels sufficiently socially supported. Finally, we asked an open ended question to specify patients’ supportive care needs by asking which form of support they wish for.

Method
Participants
We aimed at recruiting N = 300 patients. Inclusion criteria were an age of at least 16 years, the diagnosis of any rare disease, sufficient German skills, internet access and online informed consent. We excluded patients with non-rare diseases or a diagnosis that was unclear or not confirmed by a physician. Rarity was defined based on the EURORDIS definition of rare diseases (< 1:2000) [1]. If the prevalence was unavailable, a disease was included if listed in the orphan.net register of rare diseases. We further excluded participants who completed less than 85% of the survey.

We recruited between February and July 2016 via outpatient clinics, patient organizations and self-help groups for rare diseases across Germany by distributing flyers and announcing the study online and in two patient journals. Contact information of research group members was stated. After giving consent, participants anonymously completed an online questionnaire. Ethics approval was given through the ethics committee of the Hamburg Medical Counsel on February 2, 2016 (reference number PV5088).

Study design
This cross-sectional online study is part of the project ‘Patients for patients: qualified peer counselling and self-management for patients with rare chronic diseases.’ We further investigated depression and anxiety in the sample [5].

Measures
Demographic and health-related characteristics
We assessed demographic data and a variety of health-related variables. These included the diagnosis, if any family member is diagnosed with the same disease, visibility of the symptoms, comorbid diseases, current or past psychotherapeutic treatment, number of days patients have been unable to perform normal daily activities and number of doctoral visits in the last four weeks. We further asked patients if they feel sufficiently socially supported using the self-developed Item “Do you feel that you have sufficient social support?” with dichotomous response format (“yes” vs. “no”). To explore psychosocial support, we asked with whom they have supporting conversations on a regular basis. To this end, we asked the self-developed item “With whom do you have supporting conversations on a regular basis?” with the response options “nobody”, “family doctor”, “specialist physician”, “psychotherapist”, “support group”, “pastor”, “partner”, “family member”, “someone else, namely:”. All demographic and disease-related characteristics were self-reported.

Supportive care needs
We used the German short version of the Supportive Care Needs Survey (SCNS-SF34; [4]) to assess psychosocial support needs from patients’ perspective. While the instrument was developed and validated for patients with cancer [6], it has been employed investigating other patient populations [7]. The 34-item instrument measures five domains of needs (psychological, health system and information, physical and daily living, patient care and support, and sexuality needs) on a five-point response scale (1 = no need, not applicable; 2 = no need, need satisfied; 3 = low need; 4 = moderate need; 5 = high need). The psychological domain assesses needs related to emotions and coping, like dealing with anxiety. The health system and information domain assesses needs
Data analyses

We excluded patients for whom 15% or more data were missing. We calculated means and standard deviations for all metric variables as well as frequencies for all categorical variables. For categorical data, we additionally calculated 95% confidence intervals. We calculated Cronbach’s Alpha in order to determine internal consistencies of all scales and subscales used in the study. T-tests for one sample were used to compare SCNS-SF34 means in our sample with the means of a reference sample of patients with cancer, provided in the SCNS-SF34 validation study [6] based on the transformed sum scores as described above. All statistical analyses were performed using IBM SPSS 23.

Three team members evaluated answers to the open format question using qualitative content analysis [8]. We inductively derived categories from the expressed support needs. All three team members read the answers several times. One of the researchers (NU) suggested categories, which were revised by a second researcher. Based on the feedback, first drafts of the category systems were developed. The resulting first drafts of the category system were discussed with a third researcher (AW) until reaching agreement. The final category system was then used to deductively assign open format answers to categories by two researchers independently (NU, AW). The two coders compared their results and discussed discrepancies until reaching agreement. In unclear cases, a third researcher (MKD) was consulted. We used MAXQDA to support the coding process.

Results

Sample

N=304 participants with 81 rare diagnoses were included in the analysis (see Fig. 1 for flow diagram). Demographic and clinical participant characteristics are displayed in Table 1 (see Appendix Table 1, online supplementary for a list of all diagnoses). 90.5% of all participants answered the open ended question.

Social support

The majority of all participants reported not to feel sufficiently socially supported (see Table 2). Patients sought support from personal contacts, including partner, family members and self-help groups, rather than medical professionals (see Table 2). Patients report less support with increasing specialization of the medical professionals.

Supportive care needs

Standardized sum scores of the SCNS-SF34 subscales were compared to the mean values of a reference sample affected by cancer (see Fig. 2). Patients with rare diseases reported higher supportive care needs than the reference sample with respect to psychological support needs (t(283)=7.67, p<0.001 Cohen’s d=0.43), support needs regarding health system and information (t(283)=6.04, p<0.001, Cohen’s d=0.40), physical and daily living support needs (t(279)=9.15, p<0.001, Cohen’s d=0.54), supportive care needs for patient care and support (t(294)=8.47, p<0.001, Cohen’s d=0.60) and support needs concerning sexuality (t(293)=9.80, p<0.001, Cohen’s d=0.64). Overall, patients with rare disease in the current sample wished for more supportive care in all domains and their reported care needs exceeded that of the reference sample in all domains.

Supportive care needs—qualitative analysis

For elaboration, we analyzed the open-ended question. Table 3 presents the supportive care needs patients named by the subdomains listed above. In addition to these domains, patients further named the wish to be cured from their disease or reported that they currently had sufficient support. Participants asked for more and “better” information on the disease for themselves, their spouses, carers and the general public.

Better education of others who do not have the disease that they can better understand me and not label one as handicapped.
Participants reported unmet psychological needs. Qualitative analysis further revealed that, psychological needs pertain to help in developing individual strategies like disease management strategies and emotion regulation strategies. Participants reported unmet social needs in their intimate relationships. Consistent with previous research [9], participants expressed the need to be in contact with other patients with the same condition. In the present study, participants also expressed a desire for more acceptance on a societal level. This desire may stem from the experience of stigmatization. Stigma was identified as a shared challenge in a large sample of patients affected by rare diseases [10]. In a previous interview study, patients with the rare diagnosis sarcoidosis, as well as their partners, consistently reported feeling regularly misunderstood because of the general unawareness of the condition [11].

Unmet needs were further reported in the health system and information domain. Qualitative analysis further revealed that participants wished for facilitated access to experts and treatment, as well as more information about their condition that is tailored to them to aid their understanding of it. Furthermore, participants wished for additional, specific healthcare services like physiotherapy or (extended quotas of) psychotherapy. These findings are in line with a previous qualitative study, which identified topics that patients with diverging rare diagnoses

**Conclusion**

This is the first study to quantitatively and qualitatively assess supportive care needs among patients affected by rare chronic conditions. This study shows that patients with different rare diseases have high unmet support needs on different domains.
Table 1  Sociodemographic and disease-related characteristics of sample including \( N = 304 \) patients with rare diseases (\( M_{\text{age}} = 44.2, SD = 12.8, \text{range} 16–74 \))

| Variable                                           | Absolute frequency | Percentage (95% CI) |
|----------------------------------------------------|--------------------|---------------------|
| **Gender**                                         |                    |                     |
| Female                                             | 248                | 81.6 (77.2, 86.0)   |
| Male                                               | 56                 | 18.4 (14.0, 28.8)   |
| **Nationality**                                    |                    |                     |
| German                                             | 289                | 95.1 (92.7, 97.5)   |
| Other                                              | 15                 | 4.9 (2.5, 7.3)      |
| **Relationship status**                            |                    |                     |
| Single                                             | 105                | 34.5 (29.2, 39.8)   |
| Married                                            | 139                | 45.7 (40.1, 51.3)   |
| Separated                                          | 11                 | 3.6 (1.5, 5.7)      |
| Divorced                                           | 37                 | 12.2 (8.5, 15.9)    |
| Widowed                                            | 3                  | 1.0 (0.1, 2.1)      |
| Other                                              | 9                  | 3.0 (1.1, 4.9)      |
| **Living situation**                               |                    |                     |
| Alone                                              | 70                 | 23.0 (18.3, 27.7)   |
| With partner                                       | 110                | 36.2 (30.8, 41.6)   |
| Alone with children                                | 18                 | 5.9 (3.3, 8.6)      |
| With partner and children                          | 64                 | 21.1 (16.5, 25.7)   |
| With parents                                       | 24                 | 7.9 (4.9, 10.9)     |
| In institution                                     | 1                  | 0.3 (0.3, 0.9)      |
| Other                                              | 17                 | 5.6 (3.0, 8.2)      |
| **Professional status**                            |                    |                     |
| Self-employed                                      | 13                 | 4.3 (2.0, 6.6)      |
| Civil servant                                      | 6                  | 2.0 (0.4, 3.6)      |
| Employed                                           | 121                | 39.8 (34.3, 45.3)   |
| Househusband/-wife                                 | 14                 | 4.6 (2.3, 7.0)      |
| Seeking work                                       | 17                 | 5.6 (3.0, 8.2)      |
| Pension (early-age,-widow-)                        | 26                 | 8.6 (5.5, 11.8)     |
| PENSION (occupational disability/disability)       | 54                 | 17.8 (13.5, 22.1)   |
| Student                                            | 25                 | 8.2 (5.1, 11.3)     |
| Maternity/parental leave                           | 2                  | 0.7 (0.2, 1.6)      |
| Currently on sick leave                            | 4                  | 1.3 (0.0, 2.6)      |
| Other                                              | 22                 | 7.3 (4.4, 10.2)     |
| **Employment status**                              |                    |                     |
| Full-time                                          | 120                | 39.5 (34.0, 45.0)   |
| Part-time (80–99%)                                  | 20                 | 6.6 (3.8, 9.4)      |
| Part-time (60–79%)                                  | 21                 | 6.9 (4.1, 9.8)      |
| Part-time (40–59%)                                  | 18                 | 5.9 (3.3, 8.6)      |
| Part-time (< 40%)                                   | 18                 | 5.9 (3.3, 8.6)      |
| Other                                              | 107                | 35.2 (29.8, 40.6)   |
| Family members diagnosed with the same illness      | 62                 | 20.4 (15.9, 24.9)   |
| Symptoms of disease visible                        | 133                | 43.8 (38.2, 49.4)   |
| At least one self-reported comorbid disease         | 209                | 68.8 (63.6, 74.0)   |
| Psychotherapeutic treatment                        |                    |                     |
| Ever                                               | 142                | 46.7 (41.1, 52.3)   |
wished for information about, such as the impact of the disease on their daily activities. Patients in this study also wished for the information to be presented in a clear way, without foreign words that make it difficult for them to understand [9].

In the realm of physical and daily living, participants wished for more support. Responding to the open-ended question, participants also named the need for support in everyday life tasks and additionally for job-related facilitations (e.g. flexible time allocation in order to be able to attend medical appointments during work hours).

Unmet needs were reported for patient care and support. Qualitative analysis of patients’ responses showed that they wish for more sensitive communication with physicians, in particular.

Participants reported to wish for more support than they currently receive in sexual relationships in the questionnaire (sexuality needs domain), e.g. by receiving information on sexual relationships. In response to the open-ended question about support wishes, no participant named support needs with respect to sexuality. This may be because participants felt ashamed to raise these issues. Alternatively, given the open-ended format, they may have deemed the reporting of other themes more pressing. In a scoping review on patients’ supportive needs in living with ALS; loss of sexuality and intimacy was also identified as an area patients wanted help with [12].

This was the first mixed-method study to investigate supportive care needs across different diagnoses of rare conditions from the perspective of patients. A trans-diagnostic approach allows for the identification of a scope for disease management and effective care tailored to the shared needs of a large group of patients. Previous
research has primarily focused on single conditions (e.g. sarcoidosis [11]). The focus on care needs complements an emerging field of research investigating challenges faced by patients with rare diseases [10, 13]. In the face of finite healthcare resources, addressing needs of patients with rare diseases can help provide better care for a group of patients for whom healthcare professionals are otherwise confronted with the lack of a cure and knowledge about specific conditions and their management. Understanding the shared supportive care needs of groups of patients with rare diseases can therefore guide healthcare professionals in advising and treating patients with rare diseases who often present as “one of their kind” in primary care. Overall, the results of studies on single conditions are in line with the current study, despite previous studies focusing on single conditions and employing different methods for the assessment. This supports the notion that patients with different rare diseases may share generic burden and supportive care needs. Further studies on the shared needs of patients with different rare diseases qualitatively assessed the perspectives of experts on needs in the process that leads up to the diagnosis [14] or studied psychosocial supportive needs of families and care-givers [15]. The present research extends these perspectives.

A weakness lies in the sampling; it does not reflect a representative picture of patients with rare diseases. We recruited via outpatient clinics, patient organizations and self-help groups for rare diseases. It is likely that we primarily reached patients who, it can be presumed, are well integrated into care structures. Patients with rare diseases for whom there are no patient associations or specialist clinics available may have even greater support needs. More women than men participated, potentially because of their interest in the topic, which might be a source of bias. This gender ratio is similar to that of a previous study in this field on information needs [9].

**Implications and outlook: how can we improve care for patients with rare diseases?**

Assessing supportive care needs enables service providers to identify gaps in existing services and prioritize resource allocation to those aspects of care that need improvement [16]. The current study shows that adult patients affected by rare diseases need supportive care in a broad range of domains. Quality of life could be improved by implementing interventions on different levels of a patient’s ecosystem: The individual level (e.g. psychological), the patient’s microsystem (e.g. social and work environment or contact with physicians) as well as
higher order levels, i.e. the healthcare system or the general public.

Regarding the latter, patients report the need for multidisciplinary care, which could be implemented in centers for rare diseases and that in part could be coordinated through general practitioners in the absence thereof. On the level of health policy, assumption of costs for specific healthcare services, like extended quotas of

| Table 3 Support wishes according to open format answers (N = 275) |
|---------------------------------------------------------------|
| **Wishes regarding…** | **Total number of mentions** |
| **Health system and information needs** | |
| Healthcare and information | 277 |
| Facilitated access to experts and treatment | 49 |
| Information transfer for the patients | 28 |
| More support by funding agencies | 27 |
| Psychotherapy | 23 |
| Better informed physicians | 23 |
| Adequate and best possible medical care | 21 |
| Information transfer for others | 13 |
| Improved/ facilitated cooperation between different healthcare providers | 13 |
| Assistance in bureaucratic procedures | 11 |
| Guidance and consultation | 9 |
| Rehabilitation interventions | 6 |
| More research | 6 |
| Pain therapy | 4 |
| Physiotherapy | 4 |
| Support for relatives | 3 |
| Alternative treatments | 2 |
| Other treatment measures | 2 |
| Euthanasia | 2 |
| **Patient care and support needs** | |
| More personable physicians | 28 |
| Being more involved in treatment decisions | 3 |
| **Psychological Needs** | |
| Coping strategies | 4 |
| Learning disease management strategies | 3 |
| Learning emotion regulation strategies | 1 |
| Social/emotional support | 77 |
| Inclusion/more tolerance and acceptance in society | 31 |
| Social and emotional support | 22 |
| Contact to peers with the same disease/ self-help | 20 |
| Social belonging/ participation | 4 |
| **Physical and daily living needs** | |
| Support in daily life | 53 |
| Support for everyday life tasks | 25 |
| Job-related facilitations | 12 |
| Assistive technology | 11 |
| Increased disease-specific leisure opportunities | 5 |
| **Other** | |
| A symptom-free status/ cure | 8 |
| No specific support wishes | 29 |
| Uncertainty about support wishes | 12 |
| Sufficient support available | 12 |
| Unspecific support wishes | 5 |
psychotherapy could be reviewed. To further improve care for patients with rare diseases, access to experts and treatment could be facilitated by implementing telemedical measures with specialized physicians and psychotherapists. Finally, patients want more awareness of their diseases and less stigmatization, which could be achieved by information campaigns.

In the patient’s microsystem, healthcare professionals can systematically inquire needs along the dimensions reported above: whether patients feel sufficiently supported with respect to psychological needs, health system and information, physical needs, patient care, and sexuality. Based on this assessment, individual assistive measures could be taken and other providers integrated in collaborative care. Screening instruments could account for support needs and mental health, e.g. SCNS-SF34 [4] for supportive care needs, PHQ-9 for depression [17], GAD-7 for anxiety [18, 19] and SSD-12 for distressing symptom burden [20]. Furthermore, the provision of comprehensible information in written or verbatim delivery could help address patients’ information needs. For this purpose, checklists could help healthcare professionals to ensure that they have covered important issues. One way to address the need to be in contact with other patients with the same condition could be to involve expert patients in multidisciplinary care. On an individual level, implications for care can be to provide sufficient psychological aid to individually address psychological needs.

This study shows that patients with different rare diseases have high unmet support needs on different domains. Interventions addressing these needs can help to provide better and more comprehensive care. Future research could further differentiate shared needs from those specific to single conditions in order to optimize treatment.

Acknowledgements
We would like to thank all participants of the study: We thank Anne-Marie Wallmüth (AW) for assisting in qualitative analysis. We thank David Lintott for proofreading. We thank ACHSE e.V. and patient organizations that supported us in contacting their members, namely: Bundesverband Neurofibromatose e.V., Leberhilfe e.V., Pemphigus + Pemphigoid SHG e.V., Marfan Hilfe e.V., NAKOS, Marfan SHG Berlin-Brandenburg, Snyrix-Hessen e.V. (Syringomyelie Selbsthilfegruppe-Hessen-Darmstadt), Kindernetzwerk e.V. We thank all German centers for rare diseases that helped in contacting potential participants, namely: Martin Zentrum Centrums für Seltene Erkrankungen, Zentrum für seltene Erkrankungen Aachen, Zentrum für Seltene Erkrankungen – Referenzzentrum Nordbayern, Berliner Centrum für Seltene Erkrankungen, Zentrum für Seltene Erkrankungen Regensburg, Essener Zentrum für Seltene Erkrankungen (EZSE), Zentrum für Seltene Erkrankungen Mannheim, Freiburger Zentrum für Seltene Erkrankungen (FZSE), Universitäres Centrum für Seltene Erkrankungen Dresden (USE), Lübecker Zentrum für Seltene Erkrankungen, Zentrum für seltene Erkrankungen Bonn.

Authors’ contributions
Miriam Depping, Natalie Uhlenbusch and Bernd Lowe have contributed to conception, design, acquisition, analysis or interpretation of this study. Further, all authors contributed to writing or editing the draft of the manuscript and approved to the final version. Finally, all authors agree to be accountable for all aspects of the work. MKD: Conceptualization, Methodology, Formal Analysis, Investigation, Data Curation, Writing—Original Draft, Writing, Supervision, Project Administration, Funding Acquisition. Natalie Uhlenbusch: Conceptualization, Methodology, Formal Analysis, Investigation, Data Curation, Writing—Review & Editing, Visualization. Hans F. E. Klose: Writing—Review & Editing. Yskert von Kodolitsch: Writing—Review & Editing. Victor-Felix Mautner: Writing—Review & Editing. Bernd Lowe: Conceptualization, Resources, Writing—Review & Editing, Supervision, Project Administration, Funding Acquisition. All authors read and approved the final manuscript.

Funding
This study was funded by a research grant from the Robert Bosch Foundation, Germany. The study is part of research project Patients for Patients: Qualified Peer Counselling and Self-Management for Patients with Rare Chronic Diseases [ID: 32.5.A401.0004.0; Principal Investigator: Prof. Dr. Bernd Lowe, University Medical Center Hamburg Eppendorf]. The foundation provided funding but did not play a role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Availability of data and materials
Data cannot be made publically available due to ethical reasons. In the area of rare diseases, information about the diagnosis in combination with personal information may compromise anonymity and confidentiality of the participants. The independent ethics committee of the Hamburg Medical Chamber assessed our research project beforehand. The ethics vote allows sharing data with eligible researchers but we do not have approval to share the data publicly. Researchers interested in getting access to the data should feel free to contact the corresponding author (Miriam K. Depping—m.depping@uke.de) or the principal investigator (Prof. Dr. Bernd Lowe—b.loewe@uke.de). The ethics committee can be contacted at: ethik@ah-ekn.de.

Ethics approval and consent to participate
The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. Ethics approval was given through the ethics committee of the Hamburg Medical Counsel on February 2, 2016 (reference number PV5088). All participants gave informed consent and subsequently completed the online questionnaire anonymously.

Consent for publication
All authors agreed to the submission of the manuscript.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Department of Psychosomatic Medicine and Psychotherapy, University Medical Center Hamburg-Eppendorf, Martinistr. 52, 20251 Hamburg, Germany.
2 Department of Cardiology, University Medical Center Hamburg-Eppendorf, Hamburg, Germany.
3 Center for Oncology, Medical Clinic and Polyclinic, University Medical Center Hamburg-Eppendorf, Hamburg, Germany.
4 Clinic and Polyclinic for Neurology, University Medical Center Hamburg-Eppendorf, Hamburg, Germany.

Received: 18 May 2020 Accepted: 22 December 2020
Published online: 22 January 2021

References
1. Eurordis. What is a rare disease? : EURORDIS ‑ Rare Diseases Europe; 2017. [Available from: http://www.eurordis.org/sites/default/files/publications/Fact_Sheet RD.pdf.
2. Courbier S, Berjonneau E. Juggling care and daily life: the balancing act of the rare disease community. Paris: EURORDIS; 2017.
3. von der Lippe C, Diesen PS, Feraugen KB. Living with a rare disorder: a systematic review of the qualitative literature. Mol Genet Genomic Med. 2017;5(6):758–73.
4. Boyes A, Girgis A, Lecathelinais C. Brief assessment of adult cancer patients’ perceived needs: development and validation of the 34-item Supportive Care Needs Survey (SCNS-SF34). J Eval Clin Pract. 2009;15(4):602–6.

5. Uhlenbusch N, Löwe B, Härter M, Schramm C, Weiler-Normann C, Depping MK. Depression and anxiety in patients with different rare chronic diseases: a cross-sectional study. PLoS ONE. 2019;14(2):e0211343.

6. Bonevski B, Sanson-Fisher R, Girgis A, Burton L, Cook P, Boyes A, et al. Evaluation of an instrument to assess the needs of patients with cancer. Cancer. 2000;88(1):217–25.

7. Kohlmann S, Kilbert MS, Ziegler K, Schulz KH. Supportive care needs in patients with cardiovascular disorders. Patient Educ Couns. 2013;91(3):378–84.

8. Mayring P, Fenzl T. Qualitative Inhaltsanalyse [Qualitative content analysis]. In: Baur N, Blasius J, editors. Handbuch Methoden der empirischen Sozialforschung [Handbook methods of empirical social research]. Wiesbaden: Springer; 2014. p. 543–56.

9. Litzkendorf S, Babac A, Rosenfeldt D, Schauer F, Hartz T, Lührs V, et al. Information needs of people with rare diseases—what information do patients and their relatives require? J Rare Disord Diagn Ther. 2016.

10. Bryson B, Bogart K, Atwood M, Fraser K, Locke T, Pugh K, et al. Navigating the unknown: a content analysis of the unique challenges faced by adults with rare diseases. J Health Psychol. 2019. Online ahead of print.

11. Moor C, Van Manen M, van Hagen P, Miedema J, van den Toorn L, Gür-Demirel Y, et al. Needs, perceptions and education in sarcoidosis: a live interactive survey of patients and partners. Lung. 2018;196(5):569–75.

12. Oh J, Kim JA. Supportive care needs of patients with amyotrophic lateral sclerosis/motor neuron disease and their caregivers: A scoping review. J Clin Nurs. 2017;26(23–24):4129–52.

13. Uhlenbusch N, Löwe B, Depping MK. Perceived burden in dealing with different rare diseases: a qualitative focus group study. BMJ Open. 2019;9:12.

14. Blöß S, Klemann C, Rother A-K, Mehmecke S, Schumacher U, Mücke U, et al. Diagnostic needs for rare diseases and shared prediagnostic phenomena: results of a German-wide expert Delphi survey. PLoS ONE. 2017;12(2):e0172532.

15. Anderson M, Elliott EJ, Zuirynski YA. Australian families living with rare disease: experiences of diagnosis, health services use and needs for psychosocial support. Orphanet J Rare Dis. 2013;8:22.

16. Lattimore-Foot GG. Needs assessment in tertiary and secondary oncology practice: a conceptual and methodological exposition. University of Newcastle, 1996.

17. Kroenke K, Spitzer RL, Williams JB. The PHQ-9: validity of a brief depression severity measure. J Gen Intern Med. 2001;16(9):606–13.

18. Löwe B, Decker O, Muller S, Brähler E, Schellberg D, Herzog W, et al. Validation and standardization of the generalized anxiety disorder screener (GAD-7) in the general population. Med Care. 2008;46(3):266–74.

19. Spitzer RL, Kroenke K, Williams JB, Löwe B. A brief measure for assessing generalized anxiety disorder: the GAD-7. Arch Intern Med. 2006;166(10):1092–7.

20. Toussaint A, Murray AM, Voigt K, Herzog A, Gierk B, Kroenke K, et al. Development and validation of the Somatic Symptom Disorder–B Criteria Scale (SSD-12). Psychosom Med. 2016;78(1):5–12.

Publisher's Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.