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Acromegaly and the information gap: patient perceptions of the journey from 1 primary to tertiary care 2 3 Pak, Hei Yi Vivian 1,2 4 Lansdown, Andrew⁴ Taylor, Peter 3,4 5 Rees, Daffyd Aled 4,5 6 7 Davies, John Stephen ⁴ 8 Hayhurst, Caroline^{1,2} (corresponding author) 1. Department of Neurosurgery, University Hospital of Wales, Cardiff, UK 9 10 2. Cardiff University School of Medicine, Cardiff, UK 3. Thyroid Research Group, Systems Immunity Research Institute, Cardiff University 11 12 School of Medicine, Cardiff, UK 13 4. Centre for Diabetes and Endocrinology, University Hospital of Wales, Cardiff, UK 14 5. Neuroscience and Mental Health Research Institute, Cardiff University, Cardiff, UK 15 16 Name and address of corresponding author: Caroline Hayhurst, Department of Neurosurgery, University Hospital of Wales, Heath Park, 17 18 Cardiff, CF14 4XW, UK 19 Email: caroline.hayhurst@wales.nhs.uk 20 Short Title: Acromegaly and the information gap 21 Word count (excluding abstract): 3031 22 **Keywords:** Acromegaly, long-term impact, qualitative, patient perspective

Abstract: **Objective:** Acromegaly is a rare condition and there is often a long path to diagnosis for many patients. We sought to explore patient's perceptions and understanding of acromegaly, to examine the quality of communication and find gaps in the information provided at diagnosis. Design: A prospective study using qualitative research methodology and grounded theory. A semi-structured interview was conducted with 18 patients treated for acromegaly in a single tertiary centre and verbatim transcripts were thematically analysed for overarching themes. Results: 18 patients with acromegaly were interviewed. The mean age of participants was 52 (range 30 – 72). Four overarching themes emerged; (1)Patients rely on online resources to understand acromegaly in the time between diagnosis and tertiary care clinic; (2) There is not enough support available for patients; (3) Patients have a basic understanding of acromegaly and associated conditions, but the long term impact is underestimated; 4) Patients initially felt intimidated by the multidisciplinary team panel, but overall found it useful. Conclusion: Acromegalic patients have a strong need for information at the point of initial diagnosis, in particular online resources and interaction with other experienced patients. Wider dissemination of patient educational resources into primary and secondary care settings may improve overall patient satisfaction, treatment adherence and subsequent health care provider-patient relationships.

Introduction

Acromegaly, usually due to a benign pituitary adenoma, is a rare condition that has profound effects on all aspects of the body ¹. Besides causing soft tissue growth, excess growth hormone is associated with co-morbidities such as hypertension, cardiomyopathy, diabetes and cancer ². Acromegaly can be a challenging condition to identify and patients may often be diagnosed late ³. By the time they attend clinic, patients may have already developed significant health problems, with significant impact on both quality of life and life expectancy ^{4, 5}. Additionally, diagnostic delay often shapes patients' attitudes to health care professionals, which impacts on the quality of communication and subsequent treatment adherence^{6, 7}.

In UK practice, often a secondary care endocrinologist or the primary care physician conveys the diagnosis to the patient before referral to a tertiary centre multidisciplinary team (MDT). This model is increasingly the gold standard of pituitary care in North America and Europe ^{8,9}. Therefore, it is anticipated that patients are informed of their diagnosis and given background information prior to their tertiary clinic appointment to discuss treatment options, including pituitary surgery, medical treatment with somatostatin analogues or radiotherapy. However, with an incidence of only 3.8 per million population ¹⁰, many primary care and secondary care physicians may lack experience with the condition. Studies of the diagnostic pathway in acromegaly identify the point of diagnosis disclosure as critical in establishing patient- healthcare provider trust ^{6,11}. Additionally acromegaly patients are often reluctant to share concerns or questions with healthcare providers at initial consultation⁶. However, there are few studies focusing on how information should be provided and how information resources are used by patients.

Through qualitative research, this study aims to gain a better appreciation of patients' experiences with the diagnosis of acromegaly. We sought to identify gaps in patient information in the pathway

75 to treatment, in order to improve future communication and to identify the support resources 76 required and when they are needed. 77 78 79 **Patients and Methods** 80 81 Study design: 82 A prospective qualitative study using a single semi-structured interview with patients diagnosed with 83 acromegaly referred to a single regional tertiary pituitary centre. 84 85 Participants: 86 All patients aged over age 18 that had attended the pituitary multi-disciplinary pituitary clinic at the 87 University Hospital of Wales with a biochemically confirmed diagnosis of acromegaly were invited for 88 interview by telephone and provided with written information about the study. Interviews were 89 conducted via telephone or face to face. 90 91 Data collection and analysis: 92 93 Data collection and analysis in this qualitative study was based on grounded theory where ongoing 94 data analysis leads to further questions to refine evolving theories 12. As such interviews are conducted until no further themes emerge and saturation is reached¹³. This allowed systematic 95 96 generation and development of theories by being alert to emerging themes 14, prompting adding new 97 questions to the interview guide. An initial interview guide was developed and subsequent questions 98 added as new themes emerged during the study (Table 1). 99 Each interview was recorded on tape and transcribed verbatim by the interviewer. Then, the 100 transcripts were re-read to get an impression of the patients' experiences. Thematic analysis was

101 conducted first through open coding, to separate the data into segments which relate to one idea, 102 then though axial coding, to combine similar ideas into overarching themes ¹⁵. 103 Demographic data for each participant was collected including age, mode of presentation and region 104 of presentation. 105 106 Ethical considerations: 107 This study was approved by the Cardiff and Vale University Health Board Specialist Services clinical 108 board as a service evaluation and all participants provided verbal consent. 109 110 <u>Results</u> 111 25 patients were invited for interview. 18 patients were interviewed and data saturation was reached as the final interviews did not produce any new themes ¹⁶. The mean age of participants was 52 112 113 (range 30 - 72); the mean age at diagnosis was 48 (range 26 - 72). Table 2 outlines the patient 114 demographics. 11 patients (61%) were diagnosed by an endocrinologist in their local hospital, 1 115 patient received the diagnosis from an orthopaedic surgeon after referral for carpal tunnel syndrome 116 and 4 (22%) were diagnosed by their primary care physician. 117 118 Thematic analysis 119 Analysis of the interviews produced 4 overarching themes: 120 121 1) Patients rely on online resources to understand acromegaly in the time between diagnosis and tertiary 122 care clinic. 123 Since acromegaly is a rare disease, patients may be entirely unaware of the condition and hence are 124 surprised when they are given the diagnosis. Most patients felt they were not given enough 125 information at diagnosis, which is usually delivered by their local endocrinologist or general practitioner 126 (GP). Instead, they received the most information from the tertiary MDT clinic, as expected for a rare 127 disorder. Only five patients reported that printed resources would have been useful at diagnosis. 128 129 "It wasn't brilliant. There's not a real pamphlet and nothing that was explaining it. It was literally, 'you 130 got acromegaly' ... most of it was done by myself, reading online about it." 131 132 "Possibly a leaflet about... explaining things. And as I say, what the pituitary gland does and the whole... 133 some of the terminology ... of the various things that were going on." 134 135 Given delays between obtaining the diagnosis and attending the tertiary care clinic the greatest 136 information need is between appointments. All patients, except one, had searched online to learn more 137 about acromegaly. All patients who searched online did so as soon as they heard about their diagnosis. 138 However, many reported having to filter out what they read online as there is an overwhelming amount of 139 information available and some can be distressing for patients to see. 140 141 "When I googled it and there was very outdated..... there was a man with a gigantic jaw. When you 142 google things, you can get lost." 143 144 Many patients used the National Health Service website Health A-Z (www.nhs.uk) to find information on 145 acromegaly. Other resources which patients reported using are online groups such as on Facebook groups, 146 forums, videos, podcasts and blogs. To find information, patients used google with the search terms 147 'Acromegaly' or 'Pituitary Tumour'. Five patients reported they found a number of North American 148 resources online, for example, websites and video interviews, but few specific to the United Kingdom or 149 their region. Patient videos and websites from the UK may be more relevant and reassuring for the 150 patients in this study. 151

154 academic stuff from America but if I remember rightly I started looking at sort of threads and help 155 groups but it was just full of... it was the worst stories, you know, so I decided not to read about it 156 anymore. I thought it would help but actually really didn't, so I stuck with the facts, the clinical facts." 157 158 When asked if they had enough information to support treatment decisions, the majority of patients felt 159 that initially they thought surgery was the only option, but learning about the procedure and alternative 160 options helped put them at ease. With information patients felt confident in the team treatment 161 recommendation. 162 163 2) There is not enough support available for patients 164 All participants reported they would have liked to receive more support, as GPs are often unable to help 165 with issues relating to acromegaly. 166 167 "because they know you're under specialists, they can't help you because they just don't know." 168 169 Patients find the ability to contact an endocrine nurse specialist for advice reassuring but this is often only 170 available at the tertiary centre, following specialist referral. 171 172 "there is an endocrine nurse centre there. Because I know the consultant will get back to you, but he 173 very busy doing wards, doing the clinics, sometimes there is no one else that's in, to offer any advice." 174 175 Importantly, patients expressed they would have liked to talk to treated patients, to get a better sense of 176 what is happening and what they are about to go through. Meeting other people with acromegaly and 177 reaching out to support groups could be an invaluable source of support for the patients. 178

"I think the ones I remember were NHS UK, also the pituitary foundation and there seems to be a lot of

179	"Maybe meeting other people who has had it and have been cured for it so you know what they've
180	been through, what to expect. That would have been helpful."
181	
182	Earlier signposting in the primary or secondary care setting to online resources and support groups in
183	important in the patient pathway.
184	
185	"Maybe like the pituitary foundation, maybe it would have been best if I would been told to contact
186	the pituitary foundation you know? To chat with them or the pituitary nurseThat would have been
187	very good actually."
188	
189	3) Patients have a basic understanding of acromegaly and associated conditions, but the long term
190	impact is underestimated
191	
192	Despite feeling they lacked information, the majority of patients were aware acromegaly was associated
193	with a growth hormone-secreting pituitary tumour and soft-tissue growth. Almost all patients stated there
194	was not enough information given about the long-term outlook of acromegaly, but they were aware of the
195	hormone imbalance and the physical changes that would occur if left untreated. The knowledge of other
196	long-term complications of acromegaly was variable, but the majority of patients were able to name
197	several.
198	
199	"All I know it is it causes the growth hormones to grow, which umm make my fingers swell up
200	and and my feet uhh, and obviously part of my face"
201	
202	"If it's not treated, it can cause shortening of life, umm, you heart can grow and your diabetes could
203	get worse, you can have strokes, you can have a risk of heart attacks"
204	
205	"Well, the heart problems, the diabetes or the bowel problems, I don't know any more than that."

206	
207	In the beginning, patients assumed that they would be cured after the surgery and would be normal again.
208	They may not have realised that they will still require follow up for recurrence, and may still suffer from
209	symptoms after surgery, for example, headaches and fatigue.
210	
211	"It's just, I can't see an end to it, I don't feel like there will ever be an end to it. You still gotta be
212	checked after. But I can't see a light at the end of the tunnel"
213	
214	"It wasn't explained that this might not be the end of it, you might need to have another operation and
215	it might not I think I was naïve a little bit then, I hadn't realised that."
216	
217	Acromegaly had a varied long term impact on patients, 50% felt the disease had not greatly impacted their
218	lives and were able to get back to normal following surgery. However, for the remainder the experience
219	was life-changing and they still struggle with work and relationships despite successful treatment. Many
219 220	was life-changing and they still struggle with work and relationships despite successful treatment. Many patients reported ongoing problems with depression and anxiety.
220	
220 221	patients reported ongoing problems with depression and anxiety.
220221222	patients reported ongoing problems with depression and anxiety.
220221222223	patients reported ongoing problems with depression and anxiety. 4) Patients initially felt intimidated by the tertiary multidisciplinary clinic, but overall found it useful.
220221222223224	patients reported ongoing problems with depression and anxiety. 4) Patients initially felt intimidated by the tertiary multidisciplinary clinic, but overall found it useful. Patients reported being given minimal warning before attending the MDT of the nature of the clinic.
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220 221 222 223 224 225 226 227 228 229	patients reported ongoing problems with depression and anxiety. 4) Patients initially felt intimidated by the tertiary multidisciplinary clinic, but overall found it useful. Patients reported being given minimal warning before attending the MDT of the nature of the clinic. Therefore some of them felt shocked to see a large group of clinicians and nurses in the room. They were put at ease quickly and have generally benefitted from the team meeting. "It was useful I guess, you had the input of a lot of different specialists at the same time, so yeah. It was intimidating but it was also reassuring to know that there was more than one person looking at

"Well only when you go in and you see seven people sitting there and they're all, you know, professors and consultants... they were absolutely fine... but it's a bit intimidating just to see that panel..."

Patients benefit from meeting the whole team that will guide them through their treatment and follow-up, however providing information prior to the clinic on the team members and roles would be useful.

Discussion

Using qualitative research methods to explore the needs of patients newly diagnosed with acromegaly in a primary or secondary care setting demonstrates the need for high-quality, relevant online resources and local support networks. The internet is being used increasingly by patients to research their health conditions ^{17, 18}. This has been shown to affect the patient's beliefs and potentially change their decision about treatment ¹⁹. Our study shows that patients diagnosed with acromegaly turn to online resources early after initial diagnosis to learn more about their condition. Interestingly, patients appear much more reliant on online information and patient-to-patient interaction than the traditional printed information leaflet.

However, patients may be overwhelmed by what they might find online and, quality of information on the internet may be substandard ²⁰. Our cohort highlighted the importance of online materials and groups being regionally specific to them. Ideally, patients would like to be provided with reliable online resources at initial diagnosis and guidance to help them make sense of the information ²¹. In a similar study by Gurel et al ⁶, 19 patients participated in online and face-to-face interviews aimed at understanding the impact of a diagnosis of acromegaly. They demonstrated a strong desire for education about the disease at diagnosis, noting the diagnosis seemed to 'fuel a thirst for knowledge' in all participants ⁶. As in our study, participants emphasized the need for patient-patient interaction as part of their quest for knowledge and a desire to take control of their disease, highlighting the need for access to support groups. Plunkett and Barken ¹¹ suggest strategies to facilitate the patient-healthcare professional relationship throughout the

treatment pathway and highlight the provision of educational and emotional support resources at the initial diagnosis meeting. Such resources include The Pituitary Foundation (www.pituitary.org.uk). In particular, their support groups and peer support programme may address the patients' needs to speak with more experienced acromegalic patients and encourages the exchange of health information ^{18, 20, 21}. Video presentations by similar patients, such as the UK acromegaly meetup, could be helpful for newly diagnosed patients (UK Acromegaly Meetup 2017: Patient stories - Rachel and Carolyn.

https://youtu.be/llqhT-FheMA, accessed 27.03.20).

Internationally The Pituitary Society (www.pituitary-society.org), Acromunity.com and Acromegalycommunity.com provide resources and access to support groups. However, in a study to assess communication practices among endocrinologists, Polanco-Briceno et al²² reported only 14% of respondents routinely recommended educational resources or programs to patients and only 44% were aware of these resources. Additionally, most physicians in the study did not have dedicated nurse to discuss these topics with patients.

Qualitative research is becoming more popular in surgical and medical practice ²³, as it can provide an insight into the social aspects of being treated for a disease from the patient's perspective ²⁴. To date most qualitative research in acromegaly focuses on diagnostic delay and treatment adherence ^{7, 25}. A study by Sibeoni et al ⁷ conducted with 18 participants revealed the lack of awareness in the medical community as a significant factor for a diagnostic delay. Most patients had interactions with many healthcare professionals who did not recognise or believe the symptoms being reported, which in turn drives a thirst for knowledge at diagnosis and shapes subsequent attitudes towards doctors ⁶. Sibeoni et al ⁷ also reveal the psychosocial elements of a delayed diagnosis of acromegaly and suggest that endocrinologists should be involved in addressing the psychological impact of the condition together with support of mental wellbeing.

A study with a focus group of 6 acromegalic patients to explore patient perceptions of disease impact presented a wide variety of causes for a reduced quality of life in patients with acromegaly ²⁶. Some issues

discussed in the focus group correlate with those reported by patients in our study. These include fatigue, mental health problems and worries about fertility ²⁶. These issues are not covered in available disease specific quality of life questionnaires. In their study, they advise clinicians to be more aware of these problems in order that appropriate support may be provided ²⁶. It is clear that patients focus initially on the immediate treatment and resolution of symptoms such as fatigue, headaches and joint pain, but pay little attention to the potential long term implications or need for future multimodal therapy ⁶. When and how patients should receive this information and how they will process it is not clear. However our study and that of Gurel et al ⁶ suggest this is most likely to have an impact when discussed in a patient-to-patient forum.

Despite a clear benefit from a multidisciplinary team approach to the management of pituitary disease ^{9, 27}, patients find the experience intimidating. Although not all centres will adopt a joint MDT clinic with multiple clinicians; in our practice we find it beneficial to facilitate discussion between the patient, endocrinologist, surgeon and radiation oncologist to explore all treatment options. Prior contact from a specialist nurse or written information detailing the team members and their roles may help ease some anxiety and improve subsequent treatment adherence and overall satisfaction ¹¹.

Overall, despite increasing awareness of acromegaly and a recent reduction in the delay to diagnosis ⁴, most patients are diagnosed in non-specialist centres with limited access to specific support resources. There is a need for clear signposting to up-to-date online resources that patients feel is relevant to them both factually and geographically. Our study demonstrates this should occur prior to attendance at the specialist tertiary centre together with a need for supporting education on acromegaly in primary and secondary care.

Limitations:

This study involves patients from a single centre and results drawn from the study may not apply to other patient groups and/or internationally. Nevertheless, conclusions drawn from this study may still be relevant for other centres to better appreciate the needs of patients with acromegaly. In any qualitative study the interviewer's ideas and assumptions can bias the outcome; however, the semi-structured format and open questions should have allowed the patients to speak freely about their experience and concerns.

Conclusion

The study demonstrates some of the challenges faced by patients with acromegaly and gives us an insight into knowledge of their condition and what information and support they require. It highlights a need for better communication with patients as well as guidance for online searching. There is a clear need for the provision of information at the point of initial diagnosis in whatever care setting that may be made. The wider education and dissemination of appropriate online resources will improve subsequent health care provider-patient communication and ultimately improve treatment satisfaction and quality of life.

Declaration of Interest: There is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported

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