The role of gendered constructions of eating disorders in delayed help-seeking in men: a qualitative interview study

Ulla Räisänen, Kate Hunt

ABSTRACT

Objectives: To understand how young men recognise eating disorder (ED) symptoms and decide to seek help, and to examine their experiences of initial contacts with primary care.

Design: A qualitative interview study.

Setting: Men from across the UK were interviewed as part of a study of 39 young men’s and women’s experiences of having an ED.

Participants: 10 men aged 16–25 years with various EDs including anorexia nervosa and bulimia nervosa.

Results: The widespread perception of EDs as uniquely or predominantly a female problem led to an initial failure by young men to recognise their behaviours as symptoms of an ED. Many presented late in their illness trajectory when ED behaviours and symptoms were entrenched, and some felt that opportunities to recognise their illness had been missed because of others’ lack of awareness of EDs in men. In addition, the men discussed the lack of gender-appropriate information and resources for men with EDs as an additional impediment to making sense of their experiences, and some felt that health and other professionals had been slow to recognise their symptoms because they were men.

Conclusions: Although increasingly common in young men, widespread cultural constructions of EDs as a ‘women’s illness’ mean that men may fail to recognise ED symptoms until disordered behaviours become entrenched and less tractable to intervention. Men also report that such perceptions can affect the reactions of their families and friends, as well as health and educational professionals. Primary care professionals are well placed to challenge inappropriate perceptions of EDs.

INTRODUCTION

Eating disorders (EDs) are an increasingly common form of mental ill health and have been highlighted as a clinical priority area for Youth Mental Health in 2013–2016 by the UK’s Royal College of General Practitioners. Early diagnosis and intervention can prevent the development of long-term, potentially life-threatening, psychiatric morbidity with estimated costs to the National Health Service (NHS) of £50 million in inpatient care, and an additional £5–20 million for outpatient and primary care. Anorexia nervosa (AN) has the highest mortality rate of all psychiatric conditions in adolescence. Diagnostic guidelines recognise four types of EDs: AN, bulimia nervosa (BN), binge eating disorder and eating disorder not otherwise specified (EDNOS). Environmental and genetic factors impact the risk of developing an ED. Up to 70% of cases of AN and BN have been estimated to be triggered by an adverse life event, such as abuse. Existing psychiatric conditions or substance abuse in the family, certain personality traits (such as perfectionism and low self-esteem), and complex sociocultural factors around body image are also risk factors. Treatment can include psychological and pharmacological interventions as well as dietary advice, but early diagnosis and intervention is important for long-term prognosis.

Strengths and limitations of this study

- The study utilised well-developed and rigorous qualitative data collection and analysis methods.
- We succeeded in recruiting men with eating disorders, a hard-to-reach and under-researched group, who were prepared to talk openly about their experiences.
- Because these young men’s experiences were collected as part of a study that recruited and interviewed both young men and young women using the same methodology, we are able to identify what is specific to young men’s accounts of having an eating disorder.
- The study is limited to young people and does not contribute to understanding the experiences of older people with eating disorders.
EDs are commonly characterised as conditions affecting teenage girls, but there are many similarities in symptoms, treatment response, and prognosis in men and women with EDs. Although evidence on gender differences in prognosis is limited, one recent cohort study found treatment outcomes in all EDs were better for men than women, when remission was measured by weight restoration and self-reported relapsing. Some studies suggest that men with EDs are more likely to engage in obsessive exercise and have more psychiatric morbidity as a result of an ED. Homosexuality has also been cited in some studies as a risk factor for men.

Estimates of the prevalence of EDs in men and women vary widely, depending in part on whether full or partial diagnostic criteria are applied and whether studies have been community or hospital based. However, all studies suggest lower rates of EDs among men. It has been estimated that 1 in 250 women, and 1 in 2000 men in the UK have AN, and the prevalence of BN and other types of EDs is considerably higher; with BN and EDNOS thought to be more common in men than AN. A commonly cited estimate, whose provenance is unclear, is that 10% of clinical ED cases occur in men, but some data suggest that this prevalence could be as high as 25%. Despite this uncertainty about prevalence, the incidence of EDs in men is rising, although it is thought that statistics still underestimate the prevalence of EDs among men.

This is, in part, because a lack of public awareness of EDs in men may delay or prevent men from recognising early signs of EDs and seeking help. Lack of understanding and training among health professionals may also contribute to delays in diagnosing and treating EDs among men, in part because men with EDs might account for symptoms regarding weight and dieting behaviours differently from women with EDs. General practitioners (GPs) and other primary healthcare professionals play a pivotal role in recognising early symptoms of EDs and early referral for specialist care. Yet, to date, very little is known about how men recognise ED symptoms and decide to seek professional help.

Qualitative studies of men’s experiences of EDs are sparse and have focused mainly on EDs and exercise, body image, dieting practices, and sexual orientation. One recent study explored men’s experiences of recovery and another examined men’s experiences of specialist ED services. This paper presents a qualitative study of the experiences of 10 young men with EDs, whose accounts were collected within the context of a larger study of young people’s experiences of EDs. We aim to understand how young men recognise ED symptoms and decide to seek help, and their experiences of initial contacts with primary care. We focus on the following research questions:

- How do men make sense of their early (and later) signs and symptoms of disordered eating?
- How do they realise something might be wrong and require intervention?
- Are there perceived barriers to accessing primary care (or other) services for men with EDs?
- What are men’s experiences of health professionals’ responses to their initial presentations of ED signs and symptoms?

**METHODS**

**Sample and data collection**

The findings reported are part of a larger qualitative interview study with 39 young people (29 women and 10 men) with EDs in the UK, conducted to inform an online patient information resource. In this paper we focus on the experiences of the 10 men who took part (table 1).

All interviews took place in the UK, either at the interviewee’s home or a meeting room. Using maximum variation sampling, participants were recruited using a variety of channels, including patient organisations, social media, healthcare practitioners and snowballing to elicit a wide range of experiences across the illness spectrum. Inclusion criteria for the participants were: (1) aged 16–25, (2) living in the UK, (3) experience of an ED (most participants had been diagnosed but also a couple of as yet undiagnosed cases were included) and

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*Undiagnosed, but self-identified ED type(s) listed in the table. AN, anorexia nervosa; BN, bulimia nervosa; ED, eating disorder; EDNOS, eating disorder not otherwise specified.
These behaviours usually developed over months, school lunches, restrict calorie intake and eventually became habitual. The men described developing daily routines around eating and weight management, such as obsessive calorie counting, exercise and self-weighing. Their problems often also manifested in other ways, such as self-harming or increasing isolation.

Men accounted for their initial inability to attribute their feelings and behaviours to possible signs and symptoms of an ED through gendered understandings of EDs. Participant 1 ‘didn’t think for one moment [he] could be ill with an ED’ and participant 2 said the possibility of an ED ‘had not entered my head’. Participant 3 made sense of his developing routine of bingeing and purging as a personal coping mechanism rather than an indication of underlying pathology. He had never heard of BN and thought bingeing and purging was something he had ‘made up’:

I thought I made it up myself … you know, something that only I did, you know, I never thought in a million years this was something that lots of people did, and deliberately did to cause damage to themselves. You know, it wouldn’t have crossed my mind. Participant 3

Participant 4 thought of EDs as affecting ‘fragile teenage girls [who are] very emotional’. He identified himself as ‘one of the lads’ in the rugby team and not someone who could have a problem with eating. Participant 2 had always thought EDs were ‘something girls got’. Participant 1 framed his lack of awareness of EDs as congruent with society’s understanding of EDs in men:

I didn’t know men could get eating disorders then … ‘cos that would be like five years ago and there wasn’t really much said about men with eating disorders then … I didn’t know the symptoms, didn’t know anything, it was just, to me it was just happening. I didn’t really know what was going on. Participant 1

Participant 1 did not at first interpret his disordered eating behaviours as symptoms of an illness, or as anything recognisable and treatable; rather he saw his symptoms as mundane aspects of his life, something that ‘was just happening’. Indeed, all the men described having no or very little awareness of EDs in men; hence, the possibility that their symptoms could be caused by an ED, rather than reflecting idiosyncratic behaviours, was not readily considered as an explanation by men.

Men described a similar inability in friends, family and teachers to recognise their ED. Participant 5 was investigated for gastric problems for nearly a year during which time no-one, including parents and clinicians, suggested psychological reasons for his vomiting and weight loss. Participant 6 thought that people in school ‘knew’ he had a problem but it ‘was not something that was ever discussed’:

There was never any intervention by anyone [er] at school. I mean I think people knew something was wrong. I think since talking to some of my friends ...

(4) not currently receiving inpatient treatment. In-depth interviews (conducted by UR) followed a bipartite structure. Initially, interviewees were invited to talk through their experiences of having an ED in as much detail as they wished. The researcher interrupted this free narrative as little as possible. In the second part of the interview, semi-structured questions explored emerging and anticipated themes further. Questions around emerging themes provoked topics initiated by the interviewee in the early narrative and anticipated themes examined topics from existing literature. Interviews lasted between 50 min and 2 h, were audio/video recorded with consent and transcribed verbatim. Transcripts were checked against recordings for accuracy and sent to the participants for editing and approval. For research purposes all transcripts were anonymised and pseudonyms are used.

Method of analysis

Earlier analysis (by UR) of all the 39 interviews informed the development of the online patient information resource. At this stage, the 39 transcripts were closely and repeatedly read (by UR). A coding structure was developed and the data coded to anticipated and emergent themes using the Nvivo V.9 software. Based on these themes, 27 topic summaries were written and published on the http://www.healthtalkonline.org website.

This paper is based on a more detailed analysis by both authors of men’s experience of the process from symptom recognition to consultation and (potential) diagnosis. A qualitative interpretative approach was taken.38 39 Combining the methods of inductive close reading and constant comparison, subthemes in relation to the process of symptom recognition and help-seeking were examined in the accounts of the 10 men. Each coded data extract was read line by line to identify subthemes. Four subthemes were identified and analysed in more detail: (1) accounting for early signs and symptoms; (2) recognising the problem; (3) routes to help-seeking and (4) initial contact with healthcare services and information. These subthemes were summarised schematically and each occurrence of every subtheme was noted and compared with data from subsequent data extracts, using the ‘One Sheet Of Paper’ method to explore patterns and ‘deviant cases’,40 the results of which are discussed below.

RESULTS

Accounting for early signs and symptoms

All of the men took some time to recognise their experiences or behaviours as potential signs and symptoms of an ED. Those who were subsequently diagnosed with AN described how they started to skip meals, sell their school lunches, restrict calorie intake and eventually go for days without food. Those with BN described starting to comfort eat or purge after an unusually large meal. These behaviours usually developed over months,
they knew I was depressed. They didn’t know that I had an eating disorder because I don’t think any of us knew anything about eating disorders really. It wasn’t something that was ever been discussed. Participant 6

Participant 2 described family tensions when his parents suggested that he was just being ‘silly’ about his eating issues.

[They] picked up on it every now and then, like my Dad would get angry if I hadn’t had my breakfast. But it was never associated with any kind of eating disorder, ever; it was just, ‘Stop being silly’. Participant 2

Thus, a lack of awareness of EDs in men meant that unusual behaviours could be misinterpreted as personal choices. Men presented themselves as lacking the kinds of responses from their immediate surroundings which might have aided earlier recognition and help-seeking.

Recognising the problem

Symptoms and behaviours commonly continued for months or years before men began to recognise their behaviours as a problem; for many, they had become an entrenched routine. Some men described the realisation that something could be wrong as a gradual process, others as a sudden event. For some, once their problems could no longer be contained as a private ‘daily thing’ and impeded their social life or well-being, they began to recognise things were not right. Participant 7 ‘knew’ for a long time his habitual binging and purging was causing problems but had no name, nor understanding, to make sense of it. Participant 8 said his ‘personality changed’ and he became more ‘introverted’ and ‘isolated’, and participant 3 described how obsessions around eating started to ‘hold [him] back’ from ‘everything [he] wanted to do’. Participants 7 and 8 described a gradual realisation that things were ‘not normal’.

For others, initial awareness of their ED was precipitated by a crisis point that led to a hospital admission or emergency referral. These crises were often linked to marked deterioration in physical health which for many functioned as a more tangible measure of their well-being. Participant 9 described how he became ‘gaunt’, his ‘spine was protruding’ and he was growing lanugo hair. During an emergency GP home visit, participant 9 was told he required immediate hospital admission, otherwise he could be sectioned under the Mental Health Act:

He [GP] was like, ‘Either you come with me within the next twenty four hours, or I’m going to take you against your will.’ And it hit me at that point that you know, from Monday to that Tuesday when I was admitted to hospital it was a case of you know you’re working, you’re earning a living, you, you know you’re enjoying life, you know, well what, what I thought was life. But then from that Tuesday on you know, you, it’s, a complete flip side to everything. Participant 9

The trigger to recognise their ED could be precipitated by others. Participant 10 had been restricting his food intake and losing weight for a long time before his girlfriend ‘broke down’ and confronted him about his behaviour, forcing him to re-evaluate the gravity and impact of his behaviour and recognise ‘it was something I had to challenge’.

Routes to help-seeking

In addition to the delay in help-seeking attributed to a lack of self-recognition, young men discussed fears of not being taken seriously by healthcare providers or not wanting to be a ‘burden’ on services. Sometimes these fears were represented as imagined, sometimes as borne out of past experiences. Before his emergency GP referral, participant 9 had seen a GP twice with AN symptoms. He says ‘when they did see me they didn’t do anything’. He goes on to explain that the GPs had wanted to weigh him, which he refused to do in a manner typical of people with AN, but nothing then followed.

As men often recognised their problem late in their illness trajectory, when disordered behaviours were embedded, they could be hesitant to seek help because they had become reluctant to relinquish behaviours which they felt had a purpose and function in their lives, something they couldn’t ‘live without’.

It [ED] created a significant barrier for me for everything that I wanted to do or wanted to achieve. Absolutely I knew that it was creating problems at the time. It was just very difficult because you know at the same time you know I can’t live without it ‘cos I wouldn’t be able to cope, so that it was very much a conflict. Participant 3

Over time, men commonly became secretive about their behaviours, fearing judgement, forced interventions or hurting others’ feelings. This secrecy, typical of ED sufferers, could also contribute to reluctance to consult, even once people realised their behaviour was not normal. Men also described uncertainty about how and where to seek help once they became concerned about their eating:

I didn’t really know what, where to go or what to do to be honest. [Um] We’ve all heard of the like female anorexia and all of that. And everyone, I think I’d heard of anorexia [um] that isn’t what I was going through as such. And I didn’t really know what it was or where to go...

I did start googling it and I came across [um] eventually on Facebook the ‘Men have eating disorders too’, as well, and there was a couple of other websites that I looked at. [Um] But there’s still in my opinion there’s still no real information of what you do or where you go. Participant 7

Men also referred to generalised beliefs about gender-appropriate expressions of emotion and help-seeking when explaining their reluctance, as men, to seek help:
You’ve still got this again stigma attached to it [anorexia nervosa] that it’s a woman’s illness. But it’s not. And it really really annoys me because it’s not a woman’s illness, and it affects men. And I don’t know, I can’t obviously say oh [um] it’s harder for men, but there’s different experiences for men because, you know, ‘No I don’t get ill, I’m a man’. You know, ‘I don’t get ill, don’t need treatment, I don’t have emotions.’ Blah, blah, blah. Things like that. They, which could make it harder for a man to open up about it. Participant 5

Initial contact with healthcare services and information

Eight of the men had been formally diagnosed with an ED. The other two identified with prolonged ED behaviours over years but neither had yet chosen to seek diagnosis. The route to diagnosis could be complex and lengthy, even when men presented late in their illness and were in need of specialist psychiatric services. Men could face a lengthy waiting list, during which time their symptoms escalated further. Even when men had made the decision to seek help, commonly someone else, typically their mother, took them to the first appointment.

Usually the first contact with formal services was with their GP, although participant 3 preferred to see a locum GP because ‘obviously [I] didn’t have a relationship with that doctor’, so that made it a bit easier to talk about stuff and I think I was so desperate anyway I just wanted to tell somebody’ and participant 4 chose to see the school counsellor.

Once in the healthcare system, men recounted mixed experiences of interacting with health professionals. The initial appointment could dictate how willing men were to stay in touch with services. Some described positive experiences of their initial consultation. Participant 7’s GP quickly realised that he was suffering from BN, gave him information about the illness, offered a referral and wanted to monitor him but participant 7 ‘was not happy it going any further’:

I did make contact with my doctor … I just said that what I had done and what had happened… he talked through the whole process [er] with me and explained that, the very serious side effects to doing that to myself. [Um] Obviously encouraging me you know for it not to continue doing what I can. [Er] He was happy to refer me [um] to someone, but you know that was my choice. [Um] I remember at the time not wanting anything to go as far as that. And so I declined that. Participant 7

Participant 1 described his first GP contact as ‘very helpful’, acknowledging his GP’s willingness to offer support, but ultimately frustrating because:

Her [GP] struggle was that they haven’t got the services to give to you. They want to but they just haven’t got them. Participant 1

However, several men described having to go to the GP repeatedly before they felt they were being taken seriously or referred, and in a few instances an ED had been missed or misdiagnosed. Participant 2 was initially told he was ‘going through a teenage fad’ until his mother went to see the GP on her own; this precipitated a Child and Adolescent Mental Health Service (CAMHS) referral. When participant 3 first consulted and detailed his long-standing binging and purging, a locum GP actively rejected a diagnosis of BN, told participant 3 he was depressed and prescribed antidepressants, a clinical judgement which participant 3 attributed to his male gender:

At eighteen I went to the doctor, [um] I didn’t know the doctor ‘cos I was obviously new to the area, [um] and I spoke to her quite confidently about my, what I was going through, because by then I was very aware and the doctor said to me, [um], ‘You haven’t got bulimia, you’re just depressed , … I’m probably quite confident in saying that that was probably because you know I was male, you know I didn’t live up to the stereotype of being young and female, [um] and because my, it had sort of had nothing to do with body image or weight or anything like that. I think that might have confused the doctor a little bit you know. Participant 3

Instances of misdiagnoses in specialist care were also reported. Participant 5 was told by a gastroenterologist after prolonged vomiting, severe weight loss and lengthy physical investigations that his problems were not physical. Rather than referring him for psychological treatment he was, in his words, told to ‘man up’ and ‘not be weak but be strong and deal with the problem’. Participant 5 describes his reaction:

I didn’t say anything. I was dumbfounded, like I couldn’t believe a physician would say something like that, because I just thought, you know, if there’s nothing physically wrong but something’s happening surely that means there’s something psychological. And if it’s to the extent where I’ve lost three stone at this point, you know, shouldn’t I be getting referred. … basically I walked outside and punched a wall and broke my knuckle because I was so angry. That he’d sat there and called me weak and blah blah blah…

Right now it makes me so angry ‘cos there was another two weeks after that where my life went, well it didn’t go downhill but it just dropped off the edge. [Er] I wasn’t talking to anyone, I was just angry all the time, exercising more, cut myself off, and then the suicidal thoughts started coming in. And I hurt myself quite a bit, in quite a few different ways, and I eventually tried to kill myself. Whereas quite possibly if he’d referred me to even a psychiatric nurse I’m pretty certain they could have gone like that [clicked fingers], and this is what’s wrong. Participant 5

Only after an admission to accidents and emergency department following a suicide attempt were participant 5’s problems recognised and he was immediately referred to an ED clinic for intensive outpatient treatment. Men’s accounts of the readiness and capability of GPs to provide relevant and appropriate information about EDs
Men with EDs are underdiagnosed, undertreated and predominantly affecting women. Information that was made available was often viewed to be gender-blind or gender-inappropriate. Participant 1 thought the information was directed at women and described how the lack of male-specific information could provide a false sense of security for male patients. He wanted specific information on the impact of EDs on men:

like you hear like the side effects of having an eating disorder on like women. Like they can become infertile and stuff like that, but I’ve never seen any for men. So I like went and was like, ‘Look, well what are they for men?’ ‘Cos like that I could like have a side effect and I wouldn’t know.

I think I read somewhere that men can become infertile by it, and I’m like, ‘Yeah but no one tells you that’. They need to like tell you ‘this could happen, that could happen’. Like you can get, I know you can get like osteoporosis which I’ve only seen like written about women, which is obvious it could happen in men as well. So it could lead like some people to think, ‘Oh there’s no side effects for men.’ When there is and it’s just not, you just can’t, I think, I had to like scroll through the whole of the Internet trying to find bits of information. Participant 1

Only participant 5 mentioned being given male-specific information. He described the information as ‘a scare sheet’ which focused on the physical side-effects and symptoms of AN, and highlighted the possibility of impotence or lack of sex drive:

The only information I got was [er] a scare sheet basically. It was this was going to happen if you keep going. Basically the big one that they circled was, ‘Oh you won’t be able to be sexually active’ for men. And that was the biggest thing. I got about five or six different sheets from them, and basically the main fact was, ‘Oh yes, your organs won’t work, you know. You’ll lose nails, hair will thin. You cannot have sex’, big massive writing basically. Participant 5

Men expressed a strong desire for balanced, gender-tailored information. They felt that gender-blind information reflected wider societal constructions of EDs as predominately or only affecting women.

DISCUSSION

Men with EDs are underdiagnosed, undertreated and under-researched. In particular, there is a lack of qualitative research on men’s experiences of EDs and a continuing perception of EDs as a ‘female’ illness. Our findings suggest that men may experience particular problems in recognising that they may have an ED as a result of the continuing cultural construction of EDs as uniquely or predominantly a female problem. The culturally prevalent view that EDs largely affect teenage girls meant that many of these young men only recognised their behaviours and experiences as possible symptoms of an ED after a protracted delay, mitigating possibilities of early intervention and improved prognosis. Models of help-seeking recognise that, when bodily or social functioning is disrupted, people evaluate their experiences in relation to ‘illness representations’ or ‘stocks of knowledge’ that help them make sense of their symptoms and decide on appropriate actions. Thus, inappropriate understandings that EDs only affect young women can mean that men delay seeking medical advice, even when they recognise a significant deviation from the ‘healthy normal’. Wyke et al suggest that symptoms, responses and actions are simultaneously interpreted and evaluated in the light of this accumulated knowledge and through interactions with others. Thus, cultural expectations of health are an important influence on people’s own attributions of symptoms and actions, and others’ reactions. These young men’s accounts suggest their own lack of knowledge was compounded by a similar lack of recognition in their social networks. Hence, many first presented late in their illness trajectory when their ED had become well established and less tractable to treatment. This reticence could be further consolidated by broader cultural assumptions about culturally appropriate behaviours, which render young men particularly reluctant to discuss symptoms of mental or emotional ill-health. Reflecting on these findings on men in the light of the broader interview study with men and women with EDs, particular traits specific to the men’s experience seem apparent. Men and women recounted an inability in themselves and others to recognise their behaviours as symptoms of EDs but dieting and losing weight were conceptualised as more ‘normal’, unremarkable and ‘socially acceptable’ behaviours in young women. Women failed to recognise their behaviours as an ED by normalising early changes in eating and mood and discarding an ED as a ‘too severe’ diagnosis. Women did not reject an ED diagnosis as a wholly inappropriate diagnosis for them unlike men, who did not come to consider the possibility of having an ED because of the inappropriateness of an ED as a socially available explanation for them as men. Men and women recounted a reluctance to seek help which stemmed from their ambivalence towards recovery and both had experienced a prolonged diagnostic process. However, it was only men who recounted stories of explicit dismissal of their symptoms due to their gender and also reported additional challenges in lacking access to relevant and gender-appropriate information about EDs.

The study has a number of strengths and limitations. It utilised well-developed and rigorous qualitative data collection and analysis methods. Although men with EDs are a hard-to-reach group, we succeeded in recruiting men prepared to talk openly about their
experiences. The open narrative structure at the outset of the interview allowed men to spontaneously express their views about EDs as inappropriately characterised as a female-specific disease. However, since the remit of the wider study was to explore the experiences of young people with EDs; our research does not contribute to understanding the experiences of older men with EDs. This is an area that requires further research, especially in order to examine men’s developing understandings of EDs across a longer illness trajectory.

The later an ED is diagnosed, the smaller the possibility of recovery and effective therapeutic interventions, leading to increased risk of serious sequelae including mortality. Other studies note that barriers to early diagnosis include difficulty or reluctance of men to recognise their ED and a lack of confidence about how best to seek help. This is compounded by widespread assumptions that EDs are largely, or exclusively, the preserve of women, particularly teenage girls. Furthermore, gender differences in primary care consultation rates are most evident in late adolescence and early–mid adulthood; hence, GPs may have fewer opportunistic encounters in which to identify symptoms in young men.

The construction of EDs as a ‘woman’s illness’ has also been embedded in clinical practice. International Classification of Diseases (ICD) 10 and DSM IV (up to 2013) included amenorrhoea as a diagnostic criterion of AN and much patient information on EDs fails to feature men or discusses issues for women, such as the impact of food starvation on female fertility. This gender-bias might also be reinforced by current National Institute of Health and Care Excellence (NICE) clinical guidelines for the treatment and management of EDs which make no specific reference to EDs among men.

In order to improve prognosis for men with EDs, early detection is imperative. Primary care clinicians have a key role in recognising early symptoms. Until information resources are made more appropriate for men with EDs, health service providers need to be particularly sensitive to men’s needs in relation to an illness so strongly associated with women. Raising awareness of EDs more widely in society is also crucial to help men (and women) to recognise and seek help before their symptoms and behaviours become intractable. It is important to decouple the experience and (self)-management of EDs from feminised cultural imagery, resources and clinical practice if we wish to prevent men from (dis)missing signs and symptoms themselves, and prevent health and other professionals (eg, teachers) from overlooking signs and symptoms in boys and young men that they may readily recognise as indicative of EDs in young women.

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Data sharing statement Participants were invited to review their interview transcript and mark any sections they did not want used, before transferring copyright to the University of Oxford for use in research, teaching, publications and broadcasting. These carefully anonymised transcripts form part of a University of Oxford archive which is available to other bona fide research teams for secondary analysis. All researchers had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

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