When the patient is making the (wrong?) diagnosis: a biographical approach to patients consulting for presumed Lyme disease

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Background: Media coverage of Lyme disease (LD) has led to an increase in consultations for presumed LD in Europe. However, LD is confirmed in only 10%–20% of patients, with a significant number remaining in a diagnostic dead-end.

Objectives: To reach a deeper understanding of how patients themselves contribute to the diagnostic process. To describe the genesis of the LD hypothesis in care pathways.

Methods: In 2019, 30 patients from a prospective cohort consulting in the infectious diseases department at University Hospital in Marseille for presumed LD were recruited for semistructured interviews. The inclusion criteria were: suffering from subjective symptoms for 6 months, no clinical or paraclinical argument suggesting current LD. The patients’ medical trajectories were collected using a biographical approach.

Results: The diagnosis of LD was primarily triggered by identification with personal testimonies found on the Internet. Most of patients were leading their own diagnostic investigation. The majority of participants were convinced they had LD despite the lack of medical evidence and the scepticism of their referring GP.

Conclusion: GPs should first systematically explore patients’ aetiologic representations in order to improve adherence to the diagnosis especially in the management of medically unexplained symptoms. Long COVID-19 syndrome challenge offers an opportunity to promote active patient involvement in diagnosis.

Key words: clinical decision-making, Lyme disease, medical history taking, physician–patient relation, primary healthcare, qualitative research

“Cyclical movement of nature and worldly events, biology and biography: these are the two series that make life an entity at once overdetermined in its material dimension and indeterminate in its course”

Fassin D. Life: A Critical User’s Manual. Cambridge; Oxford: Polity; 2018.

Introduction

Lyme disease (LD), which received little media coverage in France until the end of the 2000s, is now the subject of controversy,1 with virulent public debates. Certain doctors and patients, represented by associations, demand recognition of a chronic form of the disease associated with nonspecific symptoms including pain, asthenia, and concentration disorders.2

The current controversies over the chronic form of LD remind us of the strength of the population’s contemporary disenchantment with science.3,4 Both the general population and the medical community are disappointed in modern science, which generates a multitude of highly specialized, fragmented, temporary, and often contradictory results, especially in the biomedical field. This is especially the case in the French context of the Lyme controversy. In 2018, French scientific societies and the National Academy of Medicine refused to approve the recommendations on LD published by the Haute Autorité de Santé,5 a French government agency. Indeed, French scientific societies (including French College of General Practitioners) did not recognize the new clinical entity (created under pressure from patient associations) called “symptom/polymorphic syndrome persisting after a possible tick bite” arguing that the term was not based on scientific evidence and opened the door to overdiagnosis and inappropriate antibiotic prescriptions.6,7

Indeed, long-term antibiotic treatments are often prescribed despite the absence of proven benefits and may cause serious adverse reactions such as catheter-related blood stream infections, pulmonary embolism, septic thrombophlebitis, and gastrointestinal bleeding, and even death.8–12

To date, there is no evidence in humans pointing towards the diagnostic criteria of a possible chronic LD.13 However, media coverage of this disease has led to an increase in consultations for presumed LD in France and in Europe.14,15 In France, annual incidence of Lyme Borreliosis is estimated at ≈33,000 cases with strong regional disparities, the incidence being very low around the Mediterranean area, where the vector is rare.16

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among patients consulting with a suspicion of LD the diagnosis is confirmed for only 10%–20%, while significant numbers of patients (6%–26%) with nonspecific symptoms (arthralgia, asthenia, myalgia, headaches) remain undiagnosed at the end of the aetiological investigation.17,18

In the context of easier access to medical information, media coverage of many health issues, and official discourse promoting patient autonomy,3 physicians are now confronted with patients who produce diagnoses and seek to confirm them through health professionals.19

This diagnostic work of the patient still constitutes an unthought/blind spot in clinical medicine as it is still taught. We consider that the lack of knowledge of how the patient produces and prioritizes diagnoses is an obstacle to the therapeutic alliance.

Using a biographical approach,20 we sought to describe the diagnostic pathways of patients who initially consulted for a suspicion of LD and for whom this diagnosis was rejected by an infectiologist at the time of their inclusion in the study. We sought to better understand firstly how patients themselves contribute to the diagnostic process; secondly to describe the genesis of the LD hypothesis in care pathways. We also wanted to better understand the role of the referring GP who in the French healthcare system has a role of coordinator of the patient’s care pathways.

Materials and methods
Study design and sampling
This qualitative single-centre study recruited participants from a regional clinical research programme dedicated to tick-borne diseases led by University Hospital Institute (IHU) Méditerranée Infections in Marseille.

Between 1 May and 30 June 2019, we recruited the first consecutive patients from a prospective cohort (n = 135) consulting in infectious diseases department at the IHU for presumed LD until data saturation was achieved. Patients were eligible to participate if they were 18 years or older, French-speaking, and accepted to sign an informed consent. Clinical and paraclinical inclusion criteria were as follows: presenting nonspecific symptoms such as fatigue, difficulty concentrating, joint, muscle, or headache pain for at least 6 months; having a negative Lyme serology (a negative western blot test); and no evidence for an organic differential diagnosis. This project was approved by an ethical committee. The reporting of this study follows the COREQ guidelines (Supplementary Figure S1).

Interview guide and biographical approach
The semistructured interviews followed a biographical approach, using an interview guide (Table 1) covering the history of symptoms and referrals to different medical specialties, detailed diagnostic pathway, including the history of the differential diagnoses, genesis of the LD diagnosis, the patient’s prioritization of the most likely diagnostic hypotheses, and the associated diagnostic degree of certainty (low, medium, and high). The interview guide also covered patients’ relationships with health professionals, the GPs role in conducting the diagnostic investigation, the impact of symptoms on daily life, and representations of the disease. The biographical approach uses a life-events calendar method (Supplementary Figure S2) consisting in a retrospective data collection tool highlighting the chronological order and proximity of events to jointly analyse several aspects of the patient’s life.20

Data collection and analysis
One male investigator (RL) trained in qualitative methods conducted all interviews. The investigator had no direct clinical relationship with any participant. Interviews occurred at the IHU after a scheduled follow-up consultation. The interviews were systematically audio recorded with the patient’s consent. Interviews lasted 63 min on average (min = 44, max = 83), The interview was completed by personal documents voluntarily provided by the patient: medical file, diary of the disease, medical diary. We also used the results of the clinical scores from the medical consultation by the infectiologist, established from validated clinical scales: the Montgomery and Asberg Depression Hetero-Assessment Scale (MADRS),21 the Fibromyalgia Rapid Screening Tool (FIRST),22 and the Fukuda diagnostic criteria for Chronic fatigue syndrome.23

Interviews were fully transcribed, coded and analysed using the NVivo qualitative data software. Two investigators (RL and CE, the clinician who performed the medical consultation) independently coded all transcripts. All the collected data were systematically cross-checked by triangulation methods. Differences were reconciled by consensus until 100% agreement was reached.

A clinical profile category was defined according to the patient’s prioritization of symptoms, in decreasing order of their impact on their quality of life. A category “diagnostic investigation coordinator” was coded from the combination of the following elements: the person who came up with the idea for the serological test, the patient’s deliberate search for a “pro-Lyme doctor” to confirm the diagnosis, spontaneous consultation of specialists (without referral by the GP), in particularly with infectious disease specialists, presence/absence of a referring GP (or other referring physician) and finally spontaneous statements during the qualitative interview such as “I conducted the investigation.” The preconstructed category “pro-Lyme doctor” was chosen if the patient reported during their healthcare pathway at least 1 consultation with a “specialist in chronic LD,” whether she/he was a doctor providing nonconventional medicines (naturopath, kinesiologist, nutrition-therapist). The preconstructed category of “Lyme activist” was attributed to the patient if he/she was a member of an association or an active member of a forum dedicated to LD.

The clinical scales were not part of the qualitative study. These clinical scales are part of a systematic diagnostic workup proposed in our centre for patients who consult for suspicion of LD. Retrospectively, we found that they were very informative for the “profile analysis” of the patients and their “main” symptoms.
Results

Participants and interview characteristics
We included 30 patients (Table 2). Participants were mainly
dominantly women (83%) with an average age of 47.3 years, with a high
education level (57% with university degree). 72% of patients
had a positive MADRS depression score, 69% had a posi-
tive FIRST test for fibromyalgia, and 76% of patients met the
diagnostic criteria for chronic fatigue syndrome.

We identified 3 main clinical profiles: the “chronic pain”
profile (included neuropathic, musculoarticular, diffuse, poorly
characterized, or headache-type pain) was predominant (50%),
followed by the “neurological” profile (27%) (included patients
whose main complaint was vertigo or sensitive motor disorders
or cognitive complaints) and the “chronic fatigue syndrome”
profile (23%) (included patients with predominant fatigue,
often associated with concentration difficulties).

The history of the disease was long with an average
symptom duration of 8.5 years (median at 5 years).

Diagnostic pathway

Recourses to secondary care physicians: numerous and
unsuccessful During their diagnostic investigation, the
patients had consulted many secondary care physicians: (often
more than 4, including a psychiatrist), they had more often
used complementary therapies (naturopathy, homeopathy,
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We send young people to psychiatry when they have Lyme disease (P14)

Perception of the GPs place in their healthcare pathway The role of referring GP as a primary care physician appeared paradoxical: although patients frequently solicited them, physicians were considered powerless, or unwilling, to offer a structured healthcare pathways.

My doctor doesn’t take my complaints seriously and won’t help me find the diagnosis anyway. (P13)
I want to be listened to (…) I want to be in good hands, I want to be directed to the right specialist, but my GP can’t deal with me (…) he doesn’t know how to handle me. (P9)
My GP got scared when we talk about Lyme or make fun of it. (P12)
My GP didn’t even take the time to read the medical record (…) I started on a good basis, it’s violent to say that it is in my head when there is a documented record! you have to respect the medical record, the work of your colleagues. (P2)
I am fed up with the diagnosis of vagal discomfort and it bothers me to always see the GP for the same thing. (P16)
It’s hard to get your doctor for a flu, so to follow a complex case. (P22)

The patient’s diagnostic work Twenty-three patients reported they had coordinated alone the aetiological investigation; for the others, the investigation was carried out by their referring GP.

I had to search by myself (…) I don’t accept that stress explains the whole picture (…) I find myself as I was at the beginning, without diagnosis, without treatment (…) I have to hold on. (P11)
I have conducted my investigation in a private clinic, they are more attentive and at least they respond to our requests! (P18)
Since I have been investigating Lyme Disease, I feel responsible for my own health (…) it’s my business after all. (P22)
I had to send my blood through a Lyme kit to Germany to confirm that I was infected with a chronic strain (…) some even have to send their blood to the vet! (P23)
I took the online Lyme diagnostic test (…) i scored very high. (P12)
I had to choose a Lyme doctor, a GP specializing in chronic lyme to be heard! (P29)
The medical record I had brought with me was quite complete! it took me a lot of time to organize it, you understand. (P16)

On the road to the right diagnosis: the feeling to be at a dead-end The long diagnostic and therapeutic wandering of these patients contributed to reinforce their feeling of being definitively misunderstood by the medical profession and that their suffering is not taken seriously. In the end, the patients’ feeling of disqualification of their own expertise explains their wandering to find a doctor who could legitimize their illness.

Table 2. Social and clinical characteristics of the 30 patients interviewed between 1 May and 30 June 2019.

| Category                        | No.          |
|---------------------------------|--------------|
| Age, mean (SD), year            | 47.3         |
| Female sex                      | 25           |
| Living in a couple              | 23           |
| Educational level               |              |
| <Secondary school education     | 7            |
| Secondary school education      | 6            |
| ≥Tertiary education             | 17           |
| Professional situation          |              |
| Active employment               | 22           |
| Unemployed                      | 3            |
| Retired                         | 4            |
| Disability                      | 1            |
| Currently on sick leave         | 17           |
| Geographical origina            | 28           |
| Provence-Alpes Côte d’Azur      | 28           |
| Clinical profile                |              |
| Chronic pain                    | 15           |
| Neurological symptoms           | 8            |
| Chronic fatigue syndrome        | 7            |
| Average duration of symptoms [min–max], year | 8.5 [0.5–54] |
| Had an average duration of symptoms ≥5 years | 15 |

aThe clinical profile was defined according to patient’s prioritization of symptoms, in decreasing order of their impact on their quality of life.

The “chronic pain” category included neuropathic, musculoarticular, diffuse, poorly characterized, or headache-type pain.

The “neurological profile” category included patients whose main complaint was vertigo or sensitive motor disorders or cognitive complaints.

The “chronic fatigue syndrome” category included patients with predominant fatigue, often associated with concentration difficulties.

kinesiology, etc.) and most of them had consulted a pain-relief centre (Table 3).

The infectiologist doesn’t take me seriously as soon as I say the word LYME, immediately closed the file, my neurologist even told me it would have been easier if I had Multiple Sclerosis. (P18)
The neurologist told me it is psychological, there is nothing more to look for. (P20).

Resistance to psychiatrization Regarding the history of differential diagnoses, most patients mentioned the diagnosis of fibromyalgia, but rejected it because they considered it as a diagnosis by default and a way of “psychiatrizing” their symptoms.

I refuse to be told that all this does not exist, that it is a figment of my imagination. (P13)
I’m tired of being told it’s all in my head or I’m having a burn out. (P2)
It’s not a comedy, it’s not fake. (P9)
Even if i appreciate the recent recognition of the diagnosis of fibromyalgia it does not explain anything (P10).
We bother the doctors to ask questions they can’t answer. (P22)
Lyme, this was a promising lead that we had started to explore and that we dropped, the doctor did not want to go any further and now we are in medical des-errancy. (P19)
Fear of being left behind diagnostically. (P1)
The constant feeling of returning to zero point. (P11)

Genesis of the LD diagnostic hypothesis
Nature of exposure to LD risk from the patient’s perspective Most patients did not report a tick bite, but often justified that they may have been bitten without noticing or remembering (Table 4). For them, the main types of potential exposure to the Lyme vector were the presence of ticks in their environment, contact with traditionally tick-carrying animals, and having spent time in a region perceived to be endemic such as forests in north-eastern France or in the Michigan (USA).

Origin of the “chronic Lyme” hypothesis The hypothesis of “chronic LD” in the diagnostic pathway/trajectory of patients was most often triggered by their identification with other patients’ testimonies circulating on different media and social networks.
I stumbled upon forums of patients who suffered from the same symptoms. One thing leading to another, they directed me to therapists who were able to listen to your history and take into consideration the human being that you are. (P2)
In this TV show, I saw myself in one of the patients who described the same pain and fatigue that no one explains while the diagnosis was obvious. (P4)
Everything leads back to Lyme when you look for information on the fibromyalgia forums, all the ‘fibros’ encourage you to have a Lyme test. (P4)
Other circumstances triggered diagnostic investigations including presence of false-positive Lyme serology during a medical check-up, family or close friends raising the question of LD and finally, this hypothesis evoked by their doctors.

Confirmation of “chronic Lyme” diagnosis All patients had previously undergone serological testing in a laboratory. Serology was often prescribed at their request, despite the fact that their referring doctors were sceptical about the Lyme hypothesis. For half of patients, the test was negative, for the other half the result was considered a false positive by the clinician according to international and national guidelines.
For patients a negative serology was not sufficient to completely exclude the diagnosis of LD and, in the case of uncertain serologies, patients often gave more weight to the positivity of the ELISA test than to a negative western blot reference test.
My doctor made it clear that he was skeptical when he agreed to prescribe me the serology (…) I was clearly advised not to get involved in Lyme serology (…) he didn’t even question it when my first test came back positive! (P17)
The first test was positive, why question it? (P12)

Some patients had used laboratories whose techniques were not validated by international standards: private laboratories in Germany or via a self-test kit obtained on the Internet, all recommended by the websites of various patient associations.
In addition, nearly half of patients had negotiated and received antibiotic therapy for “chronic Lyme disease,” which was not justified on the basis of current recommendations given the characteristics of the cases.

The issues of diagnostic work Making sense of symptoms For some patients, LD was an additional disease necessary to explain all the symptoms, when the singular clinical picture of the patient does not fit into the general framework of 1 disease. LD syndrome was clearly a way to explain all the symptoms. LD gives them the opportunity to “frame” their care pathway.
My symptoms are not typical of multiple sclerosis, especially the pain. (P26)
I gladly buy the fibromyalgia diagnosis, but I know I have something else that explains all these symptoms! (P4)
With Lyme I finally understand what is happening to me (…) and where I am going (P6)
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The majority of patients stated that they believed the diagnosis of LD to be the main explanation for their symptoms and considered this diagnosis to be highly certain.

I am convinced that I have an unrecognized chronic Lyme. (P28)
I know I have lyme because the antibiotics worked. (P23)
After searching for other diagnoses: I only see this. (P4)
For me, Lyme diagnosis is written in black and white so I don’t allow anyone to discuss it. (P2)

Denial of a severe diagnosed comorbidity LD perceived as an additional disease could sometimes play a role of a distractor: invoking Lyme rather than thinking that the symptoms were the expression of a decompensation or an aggravation of a severe chronic pathology (multiple sclerosis, heart failure, ankylosing spondylitis) for which the diagnosis had already been established and accepted by the patient.

I know deep down that I have Multiple Sclerosis but I am afraid of its evolution. I prefer to have a phony disease like Lyme. (P30)

A curable disease When patients are told all the time “it’s in your head,” the infectious aetiology was often more guilt-reducing and offers the prospect of a potentially curable. LD diagnoses were clearly perceived, as a hope for recovery.

Fibromyalgia is useless it is not curable! My rheumatologist also wants a diagnosis that we can treat! (P17)
After all these years it’s good to make people understand that it’s not our fault. (P9)
A tick bite can happen to anyone (P21)

### Table 4. Genesis of the LD diagnostic hypothesis for the 30 patients.

| Clinical and medical events                                                                 | Number of patients |
|---------------------------------------------------------------------------------------------|--------------------|
| Reported tick bite                                                                           | 8                  |
| Nature of exposure from the patient’s perspective (other than a tick bite)                   | 22                 |
|    Observed presence of ticks in their environment                                          | 6                  |
|    History of unidentified insects bites                                                     | 4                  |
|    Contact with traditionally tick-carrying animals                                          | 5                  |
|    Tick-bite episode in the entourage                                                       | 2                  |
|    Endemic region                                                                            | 5                  |
| Confirmed history of erythema migrans                                                       | 3                  |
| Origin of the “chronic Lyme” hypothesis                                                     |                    |
|    Identification with clinical narratives (TV, media, Internet)                             | 14                 |
|    Physician                                                                                | 5                  |
|    Entourage                                                                                | 5                  |
|    Medical check-up                                                                         | 6                  |
| Lyme serology performed in private laboratories                                             | 30                 |
| Results of Lyme serology test                                                               |                    |
|    Negative                                                                                 | 16                 |
|    False positive                                                                           | 14                 |
| Serology performed in a nonapproved laboratory                                              | 8                  |
| Internet diagnostic self-questionnaire                                                      | 15                 |
| Received “anti-chronic Lyme disease” antibiotic treatment                                    | 14                 |
| Pro-Lyme Doctor intervention during their diagnostic pathway                                 | 8                  |
| Members of a pro-Lyme association ("Lyme disease activists")                               | 3                  |
| Have requested and obtained a doctor’s prescription for an LD serological test               | 17                 |
| Referring physician’s position on the Lyme hypothesis                                        |                    |
|    Proactive                                                                                | 5                  |
|    Neutral                                                                                  | 14                 |
|    Sceptical                                                                                | 9                  |
|    Absent                                                                                   | 2                  |
| Patient’s diagnostic hypotheses ranking                                                     |                    |
|    LD hypothesis rank first                                                                  | 24                 |
| The degree of certainty associated with the diagnoses among patients ranking Lyme hypothesis first |          |
|    High degree of certainty                                                                 | 13                 |
|    Moderate–low degree of certainty                                                          | 11                 |
Discussion

When patients make the diagnosis

Our study shows important role of narratives from other patients on social networks or in the media. These narratives are particularly valued by patients in situations marked by their perception of the lack of satisfactory diagnostic proposal from doctors. The general public refers to this type of information source much more often, to the detriment of more “objective” and official sources. This is in line with the results of a qualitative survey performed in Connecticut (USA), which reported that patients with LD placed greater trust in the experiences of close relatives who had contracted LD than in information disseminated by health professionals and health authorities. Numerous studies indicate personal testimonials play an important part on how people form opinions about health issues such as vaccination.

Most of patients managed to convince their GP to prescribe a Lyme serology test, illustrating that the medical decision is no longer monopolized by doctors. This reflects the contemporary role of patients claiming the legitimacy of a diagnosis based on their own experience. Studies recently described that diagnostic work was more particularly exercised by patients when physicians are unable to elucidate the causes of their disorders than when they are, with patients taking charge of the entire sequence from self-diagnosis to self-prescribing. Moreover, patients can nowadays find online tools (diagnostic self-questionnaire) promoted by patient groups to back up their hypotheses. Finally, self-initiated diagnostic investigation increased the degree of belief in the diagnosis of Lyme.

A diagnosis set in advance

One of the striking findings of our study is the high degree of belief in the diagnosis of LD, despite the absence of objective biological evidence usually accepted by the medical profession. The “Lyme activist” patient profile, and/or an encounter with a “pro-Lyme doctor” concerned a minority of the healthcare pathways described in this study and cannot by itself explain this high level of conviction.

The attribution of symptoms to a well-identified external cause (a bacterium) is frequent in the literature on LD and more generally on somatoform disorders. The infectious origin is often more guilt-reducing for patients and offers the prospect of a possible cure. Even though these patients may have mental health problems, they experienced the psychiatric framing as a form of violence.

Moreover, the higher level of certainty about the LD hypothesis in patients leading their own diagnostic pathway suggests they had a preestablished aetiological scenario and were seeking to put together the different elements of the medical puzzle to support it. In cognitive psychology, this phenomenon is known as “confirmation bias”: human beings tend to seek, value, and recall information that confirms or supports their previous personal beliefs.

The patients’ reasoning included 2 additional cognitive biases: firstly, a groupthink bias which allows patients with somatoform disorders to access a positive identity and thus reduce the social pain of feeling rejected by the medical profession or having the legitimacy of their illness questioned; secondly a hidden cost bias which locks patients into a misdiagnosis because of the considerable amount of time, effort, and resources already invested in the LD diagnosis and the difficulty of reinvesting in an alternative hypothesis.

Such biases clearly exacerbate false belief entrapment and isolation of the patient in the care pathway.

Strengths and limitations

The originality of this study lies in the population studied which consists of patients who have reached a diagnostic dead-end. There are very few studies on the care pathways of such patients.

To our knowledge, this study was also the first to apply a biographical approach to the diagnostic trajectories of patients consulting in infectious disease wards for a suspected LD. It allowed for the joint analysis of contextualized self-reported data and clinical data from medical records. The interviews, by focussing on the overlap between life, medical, and clinical events, highlighted the 3 dimensions at work in any care pathway: biology, biography, and social context.

As for any qualitative study, it is important to be cautious about generalizing these results. In addition, convenient sampling method introduces motivation bias into the study in particular to over-select patients who are more reflective and engaged in their care pathway. However our sample was relatively diversified with a patient profile quite similar to the rest of the cohort of patients coming to consult for presumed LD. Patients with a “Lyme activist” profile were marginal in our sample. In addition our sample size (n = 30) was the largest to date among the international qualitative research published on the subject.

Comparison with existing literature

These sociodemographical characteristics of patients (tend to be female, of middle age and have more education) are similar to those of population with chronic fatigue syndrome attributed by patients to a virus in the in the 1990s and 2000s. They are also similar to those of the Complementary and Alternative Medicines (CAM) users: 18 out of 30 patients use CAM in the present study.

A previous qualitative study involving 13 patients in the Savoy region of France reported similar results regarding the role of the Internet and the media in patients’ care pathways and in triggering their suspicion of chronic LD. However, the method of recruitment through a patient association led to an overrepresentation of “activist” patients, who were more likely to support conspiracy theories, who were explicitly reported to be in conflict with the medical profession, and who had been “exposed to Lyme doctors.” Other international qualitative studies on the subject focussed only on the experience and impact of the disease in patients’ daily life.

Conclusions

GP’s role proposes concrete solutions to improve the quality of life of these patients. With LD, the stakes have shifted to the diagnostic field. The clinician’s ability to listen to patients’ disease history rarely includes consideration of their diagnostic experience. Opposing the doctor as the sole custodian of the medical diagnosis, to patients reduced to the subjectivity of their symptoms, runs the risk of diagnostic dead-ends or parallel diagnostic pathways. Dissatisfaction with the
medical diagnosis is the classic explanation for the use of alternative medicine.\textsuperscript{43,44} Finally, the empowerment of patients in the diagnostic process suggests that doctors in France in particular are insufficiently trained to deal with functional somatic syndrome.\textsuperscript{25}

“Inductive foraging”\textsuperscript{30} could also be a relevant strategy to involve patient in the diagnostic process: at the beginning of the consultations, allow time for the patient to freely reveal the reasons for consultation and the diagnoses they think of. While the patients actively present their complaints, the general practitioners listen and take up the elements of their presentation. Faced with erratic beliefs, the reaffirmation of an assertive professional posture seems necessary: demonstrate professional expertise in a respectful manner without humiliating or demeaning the patient, while respecting the patient’s right to decide for themselves.\textsuperscript{47} We think this patient-centred care approach is likely to improve adherence to the functional somatic syndrome diagnosis and the therapeutic alliance. We are probably already facing a similar challenge with the recognition of patients with subjective symptoms in the context of a Long COVID for whom studies show a significant alteration in quality of life.\textsuperscript{46,47}

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Supplementary material

Supplementary material is available at Family Practice online.

Ethical approval

The study was conducted in accordance with the Declaration of Helsinki and approved by was approved by an ethical committee (French Committee for the Protection of Persons, authorization no. 2019 T3-10). Written informed consent was obtained from all subjects involved in the study.

Conflict of interest

The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

Data availability

The data presented in this study are available upon request from the corresponding author.

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