A fever of unknown origin as a presenting symptom in toxoplasmosis: Back to earth

Jill F. Mentink, Michiel van Rijn, Adriaan Dees

ABSTRACT

Introduction: Fever of unknown origin (FUO) and weakness in the upper legs are among the presenting symptoms in acute toxoplasmosis, which can occur in immune-competent individuals. Case Report: A 76-year-old Caucasian male presented at the hospital with relapsing fever and weakness in both upper legs. A neurological examination confirmed weakness in the quadriceps muscles, which appeared to be atypical for any neurological substrate. An extensive work-up did not lead to a diagnosis. After two weeks of clinical investigations and persistent fever, we re-examined our patient to look for anamnestic signs. We again asked about (recent) vacations, hobbies and potential exposures. Ultimately, the patient told us he performed charity work at a care farm for more than ten years. He had contact with sheep and soil, which occurred while working in the garden without wearing gloves. A diagnosis of a primo toxoplasmosis infection was made and the patient was treated with dapsone and clindamycin. Within two days his temperature normalized and the strength in his legs improved gradually. He achieved a complete recovery within three months. Afterwards his wife mentioned that her husband was afraid of losing his volunteer job, when telling his doctors about his work with disabled people. Conclusion: In patients presenting with FUO, clinicians in search of a diagnosis must struggle through a long list of possible diagnosis. Diagnostic delay can be shortened by considering the potential clues in the patient’s history.
A fever of unknown origin as a presenting symptom in toxoplasmosis: Back to earth

Jill F. Mentink, Michiel van Rijn, Adriaan Dees

ABSTRACT

Introduction: Fever of unknown origin (FUO) and weakness in the upper legs are among the presenting symptoms in acute toxoplasmosis, which can occur in immune-competent individuals. Case Report: A 76-year-old Caucasian male presented at the hospital with relapsing fever and weakness in both upper legs. A neurological examination confirmed weakness in the quadriceps muscles, which appeared to be atypical for any neurological substrate. An extensive work-up did not lead to a diagnosis. After two weeks of clinical investigations and persistent fever, we re-examined our patient to look for anamnestic signs. We again asked about (recent) vacations, hobbies and potential exposures. Ultimately, the patient told us he performed charity work at a care farm for more than ten years. He had contact with sheep and soil, which occurred while working in the garden without wearing gloves. A diagnosis of a primo toxoplasmosis infection was made and the patient was treated with dapsone and clindamycin. Within two days his temperature normalized and the strength in his legs improved gradually. He achieved a complete recovery within three months. Afterwards his wife mentioned that her husband was afraid of losing his volunteer job, when telling his doctors about his work with disabled people. Conclusion: In patients presenting with FUO, clinicians in search of a diagnosis must struggle through a long list of possible diagnosis. Diagnostic delay can be shortened by considering the potential clues in the patient's history.

Keywords: Diagnostic delay, Fever of unknown origin (FUO), Toxoplasma gondii

INTRODUCTION

Fever of unknown origin (FUO) and weakness in the upper legs are among the presenting symptoms in acute toxoplasmosis, which can occur in immune-competent individuals.

The prevalence of the obligate, intracellular protozoan Toxoplasma gondii varies worldwide, but it is generally assumed that the global infection rate is 25–30%. Some healthy individuals who are infected become
symptomatic. Immunocompromised patients are at risk of reactivation of a primo infection and developing a severe and complicated illness.

In patients presenting with FUO, clinicians in search of a diagnosis must struggle through a long list of possible diagnosis. In this modern era of sophisticated imaging techniques, the temptation to fully rely upon laboratory tests results and scans should be resisted [1, 2]. This diagnostic delay can be shortened by considering the potentially diagnostic clues in the patient’s history.

CASE REPORT

A 76-year-old male was admitted to hospital with relapsing fever along with weakness and pain in his upper legs. He said told that the weakness, which had a progressive course, had begun several weeks earlier and had progressively worsened. His medical history included diabetes mellitus type 2, hypertension, uncomplicated coronary angiographic bypass graft and laparoscopically performed cholecystectomy. Moreover, 24 years earlier, he had undergone allogeneic bone marrow transplantation because of paroxysmal nocturnal hemoglobinuria (PNH). His daily medication regime included metformin, enalapril, fluvastatin and acetylsalicylic acid.

On admission the patient was in a poor condition. He had lost 4 kg, was anorectic and had a temperature of 39.0°C. On physical examination, his heart and lungs were normal. Neurologic examination: the patient was alert, febrile, Glasgow Coma Scale E4M6V5. Investigation of the lower extremities: the patient was unable to lift his feet, the sensibility was normal. Knee and Achilles reflexes were 0/0 and -4/-2 respectively.

Laboratory tests on admission: leucocytosis of 15.9x10^9/L, sedimentation rate in the first hour of 90 and C-reactive protein of 386 mg/L. Blood and urine samples were obtained for cultures, full blood count and chemistry, along with serological tests for underlying infectious causes and rheumatic diseases. A chest X-ray and abdominal ultrasound were performed. A presumptive diagnosis of infection was made and because of the lack of focus we decided not to start antibiotics but wait for the results of the cultures. A neurologist was consulted, but found no central or peripheral abnormalities, apart from painful quadriceps muscles, that might explain the clinical picture. We did not perform a muscle biopsy at that moment. During his stay in the hospital, a relapsing fever was observed, that met the criteria of FUO, also known as febris e causa ignota (Figure 1). Cultures of blood and urine remained negative. Autoimmune serology and ANCA’s were absent. Based on serology hepatitis, HIV, Borrelia, influenza, syphilis, Coxiella burnetii (Q-fever) and brucellosis were ruled out. The serological tests for Epstein—Barr and cytomegalic virus were positive for past infections. Withdrawal of fluvastatin did not result in any improvement. A thoracic and abdominal computer tomography (CT) scan did not reveal solid masses, abscesses, lymphadenopathy or any sign of malignancy, except for a benign cyst in the liver. A whole body (18) F-fluorodeoxyglucose ((18)F-FDG) position emission tomography (PET)-CT scan showed some, perhaps physiological uptake in the kidneys and bowel. Bone marrow aspiration and immunology analyses revealed normal cell lines following the previous transplantation, without signs of leukaemia or myeloproliferative disease. Throughout this diagnostic process, we failed to arrive at a definite diagnosis. The patient remained febrile and his condition progressively worsened in a couple of days. We decided to return to the patient to seek out potentially diagnostic clues that we had previously neglected. Subsequently, the patient mentioned that he had travelled to Brazil several years ago, but was not sure whether he had received all the proper vaccinations. Thereafter, a thick blood smear was performed to exclude malaria parasites, as well as tests for schistosomiasis and Q-fever. Eventually, the patient told us that he had worked as a charity worker for more than ten years at a care farm for disabled people. He had contact with sheep and also maintained the gardens without wearing gloves while working with soil. Afterwards his wife told us that he had been very afraid to lose his volunteer job, as a consequence of telling his doctors this information.

Finally, we found that toxoplasmosis serology for IgM and IgG were positive with a rather low avidity of 0.548, which fit the diagnosis of acute infection. We considered this a primo infection in an immunocompetent host.

We started our patient on daraprim (pyrimethamine) and clindamycin. Daraprim is a folic acid antagonist that is highly effective against toxoplasma especially when used in combination with sulphonamide. We preferred the combination with clindamycin because of less symptoms and similar effectiveness. We started with one dose of 100 mg Daraprim and continued with 50 mg once daily in combination with clindamycin 600 mg four times daily for two weeks.

---

Figure 1: Temperature curve.
The patient was regularly seen at outpatient clinic after his discharge from hospital. His condition gradually improved, without a return of complaints or fever. The patient remained seropositive (IgM and IgG) during six months of follow-up.

DISCUSSION

This report illustrates the pitfalls in the diagnostic route of a patient, who may present with a fever of unknown origin. During the ward grand rounds we discussed the immune status of this patient. He was known to have diabetes type two. He exhibited no previous signs of recurrent infections. He had undergone bone marrow transplantation because of a benign disease more than 20 years ago. However, after we made the diagnosis, we wondered whether our initial assumption was right. Very few healthy individuals who have been infected with *Toxoplasma gondii* become symptomatic, although fever and myalgia have been described as presenting symptoms, even in immunocompetent individuals [3, 4].

The IgG and IgM antibodies were positive at presentation, with a low avidity of 0.548 for IgG. The IgM antibodies usually present within the first week of presentation, peak in one month, but may persist for a year or more. The IgG titer helps to distinguish an acute infection from a past infection [5]. A low avidity is associated with an acute infection, a high avidity makes an past infection or re-activation more likely.

*Toxoplasma gondii* is an obligate intracellular protozoan. The prevalence of this single cell parasite differs worldwide, but is generally assumed that that the infection rate of the global population is 25–30% [6]. A high prevalence infection occurs in tropical Africa and Latin America. A total of three major genotypes have been identified, among which type I is the most virulent. Cats facilitate the sexual parasitic cycle and spreading of oocysts through faeces. Humans can acquire *Toxoplasma* by eating undercooked meat, drinking contaminated water, swallowing parasites after contact with cat feces (via contaminated soil or cleaning a cat litter box) or they are infected by receiving an infected organ transplant or donor blood. Some infected individuals experience non-specific flu-like symptoms. By contrast, immunocompromised patients are at risk of developing severe toxoplasmosis infections after primary infection or re-activation that may result in myocarditis, pneumonitis, polymyositis, meningoencephalitis or other conditions. Morbidity is even higher after infection with a type one or atypical strain. Afterwards we asked the transplantation centre that had previously treated the patient, whether his toxoplasmosis serology had been noted in the charts prior to transplantation. This information could not be located.

The role of the FDG-PET/CT scan may be questioned. This nuclear imaging test has shown its utility in patients with FUO, along with reasonable performance [7].

In the patient presented here, the diagnostic yield of CT and PET was unclear. In contrast to the imaging techniques, our search for specific diseases based on clinical reasoning and serology was more successful [8].

CONCLUSION

The case demonstrates that FUO and weakness in the upper legs can be the presenting symptoms in toxoplasmosis and that this might present in immune-competent individuals. Unravelling the patient history can shorten the diagnostic delay. The diagnostic clue in this case turned out to be the patient’s previous social work on a charity farm. The clinical course of the patient suggested an acute infection, however a reactivation from a prior infection, in case of a transplantation history, could not be excluded definitely.

Author Contributions
Jill F. Mentink – Substantial contributions to conception and design, Acquisition of data, Analysis and interpretation of data, Drafting the article, Revising it critically for important intellectual content, Final approval of the version to be published
Michiel van Rijn – Analysis and interpretation of data, Drafting the article, Revising it critically for important intellectual content, Final approval of the version to be published
Adriaan Dees – Substantial contributions to conception and design, Acquisition of data, Analysis and interpretation of data, Drafting the article, Revising it critically for important intellectual content, Final approval of the version to be published

Guarantor
The corresponding author is the guarantor of submission.

Conflict of Interest
Authors declare no conflict of interest.

Copyright
© 2017 Jill F. Mentink et al. This article is distributed under the terms of Creative Commons Attribution License which permits unrestricted use, distribution and reproduction in any medium provided the original author(s) and original publisher are properly credited. Please see the copyright policy on the journal website for more information.

REFERENCES
1. Brown M. Pyrexia of unknown origin 90 years on: A paradigm of modern clinical medicine. Postgrad Med J 2015 Dec;91(1082):665–9.
2. Wei HX, Wei SS, Lindsay DS, Peng HJ. A Systematic Review and Meta-Analysis of the Efficacy of Anti-Toxoplasma gondii Medicines in Humans. PLoS One 2015 Sep 22;10(9):e0138204.
3. Cuomo G, D’Abrosca V, Rizzo V, et al. Severe polymyositis due to Toxoplasma gondii in an adult immunocompetent patient: A case report and review of the literature. Infection 2013 Aug;41(4):859–62.
4. Abhilash KP, Roshine MK, Vandana K, Varghese GM. A probable case of acquired toxoplasmosis presenting as pyrexia of unknown origin in an immunocompetent individual. Int J Infect Dis 2013 Nov;17(11):e1067–8.
5. Lappalainen M, Hedman K. Serodiagnosis of toxoplasmosis. The impact of measurement of IgG avidity. Ann Ist Super Sanita 2004;40(1):81–8.
6. Robert-Gangneux F, Dardé ML. Epidemiology of and diagnostic strategies for toxoplasmosis. Clin Microbiol Rev 2012 Apr;25(2):264–96.
7. Takeuchi M, Dahabreh IJ, Nihashi T, Iwata M, Varghese GM, Terasawa T. Nuclear Imaging for Classic Fever of Unknown Origin: Meta-Analysis. J Nucl Med 2016 Jun 23. pii: jnumed.116.174391.
8. Martino R, Maertens J, Bretagne S, et al. Toxoplasmosis after hematopoietic stem cell transplantation. Clin Infect Dis 2000 Nov;31(5):1188–95.

Access full text article on other devices

Access PDF of article on other devices
Edorium Journals: An introduction

Edorium Journals Team

About Edorium Journals
Edorium Journals is a publisher of high-quality, open access, international scholarly journals covering subjects in basic sciences and clinical specialties and subspecialties.

Invitation for article submission
We sincerely invite you to submit your valuable research for publication to Edorium Journals.

But why should you publish with Edorium Journals?
In less than 10 words - we give you what no one does.

Vision of being the best
We have the vision of making our journals the best and the most authoritative journals in their respective specialties. We are working towards this goal every day of every week of every month of every year.

Exceptional services
We care for you, your work and your time. Our efficient, personalized and courteous services are a testimony to this.

Editorial Review
All manuscripts submitted to Edorium Journals undergo pre-processing review, first editorial review, peer review, second editorial review and finally third editorial review.

Peer Review
All manuscripts submitted to Edorium Journals undergo anonymous, double-blind, external peer review.

Early View version
Early View version of your manuscript will be published in the journal within 72 hours of final acceptance.

Manuscript status
From submission to publication of your article you will get regular updates (minimum six times) about status of your manuscripts directly in your email.

Our Commitment

Six weeks
You will get first decision on your manuscript within six weeks (42 days) of submission. If we fail to honor this by even one day, we will publish your manuscript free of charge.*

Four weeks
After we receive page proofs, your manuscript will be published in the journal within four weeks (31 days). If we fail to honor this by even one day, we will publish your manuscript free of charge and refund you the full article publication charges you paid for your manuscript.*

Favored Author program
One email is all it takes to become our favored author. You will not only get fee waivers but also get information and insights about scholarly publishing.

Institutional Membership program
Join our Institutional Memberships program and help scholars from your institute make their research accessible to all and save thousands of dollars in fees make their research accessible to all.

Our presence
We have some of the best designed publication formats. Our websites are very user friendly and enable you to do your work very easily with no hassle.

Something more...
We request you to have a look at our website to know more about us and our services.

* Terms and condition apply. Please see Edorium Journals website for more information.