The Financial Implications of a Well-Hidden and Ignored Chronic Lyme Disease Pandemic

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Abstract: 1 million people are predicted to get infected with Lyme disease in the USA in 2018. Given the same incidence rate of Lyme disease in Europe as in the USA, then 2.4 million people will get infected with Lyme disease in Europe in 2018. In the USA by 2050, 55.7 million people (12% of the population) will have been infected with Lyme disease. In Europe by 2050, 134.9 million people (17% of the population) will have been infected with Lyme disease. Most of these infections will, unfortunately, become chronic. The estimated treatment cost for acute and chronic Lyme disease for 2018 for the USA is somewhere between 4.8 billion USD and 9.6 billion USD and for Europe somewhere between 10.1 billion EUR and 20.1 billion EUR. If governments do not finance IV treatment with antibiotics for chronic Lyme disease, then the estimated government cost for chronic Lyme disease for 2018 for the USA is 10.1 billion USD and in Europe 20.1 billion EUR. If governments in the USA and Europe want to minimize future costs and maximize future revenues, then they should pay for IV antibiotic treatment up to a year even if the estimated cure rate is as low as 25%. The cost for governments of having chronic Lyme patients sick in perpetuity is very large.

Keywords: Borrelia; Lyme disease; chronic Lyme disease; ILADS; incidence rate; cost chronic Lyme disease; CDC

1. Introduction

The objectives of this article are to investigate the incidence rate of Lyme disease in the USA and Europe, to investigate the financial cost of chronic Lyme disease and to find the most cost-efficient way for the governments to solve the current chronic Lyme disease pandemic [1]. Very few studies exist that calculate the economic impact of chronic Lyme disease [2]. According to the European Centre for Disease Prevention and Control (ECDC), Lyme disease—also known as Borreliosis—is caused by a spirochete bacteria called Borrelia burgdorferi [3]. An early stage Borrelia infection is known as acute Lyme disease and a late stage Borrelia infection is known as chronic Lyme disease. A Lyme disease “war” has been going for a long time [4,5] between two doctors’ associations regarding the most appropriate way to diagnose and treat Lyme disease. The two doctors associations are the Infectious Diseases Society of America (IDSA) [6] and International Lyme and Associated Diseases Society (ILADS) [7]. IDSA represents infectious disease doctors that firmly believe that all Lyme disease infections, regardless of whether the infection is acute or chronic, can successfully and easily be treated with three weeks of oral antibiotic [8] despite the fact that no scientific studies currently exist that support the claim that three weeks of oral antibiotics can always cure acute or chronic Lyme disease [9] and there exist many scientific studies that have shown that the Borrelia bacteria can survive three weeks of oral antibiotics in vitro [10,11] and in vivo, in mice [12–15], dogs [16], horses [17], monkeys [18] and in humans [19–36]. The in vivo references have mostly been extracted from [37–39]. Alive Borrelia bacteria has been found in 7 out of 8 patients with chronic Lyme disease [40]. The immune system by itself can never eradicate a Borrelia infection [41–44] and no scientific studies exist that show that
bacterial infections can simply disappear. Approximately 63% [45,46] of people today that are infected with *Borrelia*, unfortunately, develop chronic Lyme disease. Most Lyme disease patients today either do not receive any antibiotic treatment at all or they receive an insufficient amount of antibiotic treatment. The IDSA does not recognize chronic Lyme disease, nor do they recommend antibiotic treatment for chronic Lyme disease. The IDSA calls chronic Lyme disease post-treatment Lyme disease syndrome (PTLDS) [47] but it has been suggested that this term is a misnomer and should not be used [48]. Because chronic Lyme disease is classified as a syndrome instead of an infection, the IDSA therefore does not recommend antibiotic treatment for chronic Lyme disease.

In 2006, the IDSA treatment guidelines development process was subjected to an antitrust investigation by the Connecticut Attorney General [49], which found that many IDSA treatment panel members had conflicts of interest [50]. In November 2017, another antitrust lawsuit (Civil Action No 17-cv-190) [51,52] was directed at the IDSA along with several large insurance companies in the USA and several IDSA medical doctors, in a federal court in Texarkana, Texas in the USA. The allegations concern violations of the Racketeer Influenced and Corrupt Organizations (RICO) Act and the Sherman Antitrust Act. According to the Agency for Healthcare Research and Quality (AHRQ), which is part of the U.S. Department of Health and Human Services, 86% of health care costs in 2010 in the USA came from patients with one or more chronic conditions [53]. The concern is that because Intravenous (IV) antibiotic treatment for people with chronic Lyme disease costs so much money, insurance companies in the USA may gain by colluding with the IDSA to ensure chronic Lyme disease is not classified as a real disease caused by bacteria, so that IV antibiotic treatment can be denied.

The scientific process fails if it can be controlled by a group or groups with a particular agenda [54]. The IDSA’s treatment guidelines for Lyme disease are problematic in the following ways: (A) They do not follow The National Academies of Sciences, Engineering and Medicine recommendations on how to develop treatment guidelines [55] because the IDSA treatment guideline panel did not consist of a diversified and balanced group of doctors nor did the panel include patient representatives; (B) More than 50% of IDSAs treatment guidelines are based only on “expert opinion” [56]; (C) The IDSA’s treatment guidelines for Lyme disease do reflect the fact that, according to the CDC, Lyme disease can lead to death if not treated with antibiotics [57]. The CDC has, however, significantly underestimated the number of deaths caused by *chronic* Lyme disease. One study estimates that chronic Lyme disease and associated diseases could be the cause of over 1200 suicides per year in the USA [58].

The International Lyme and Associated Diseases Society (ILADS) represents Lyme Literate Medical Doctors (LLMDs) from around the world who recognize that chronic Lyme disease is a real and serious disease that must be treated with long-term antibiotics [59]. ILADS’ treatment guidelines for Lyme disease can be found on the National Guidelines Clearinghouse (NGC) web page [60]. According to ILADS, Lyme disease is a clinical diagnosis that is based on the symptoms a patient presents. LLMDs prefer to treat chronic Lyme disease patients with IV antibiotics because IV antibiotics are more effective than oral antibiotics [61]. However, oral antibiotics are much easier to administer, and they are not as expensive as IV antibiotics. Hence, in a situation where a patient cannot afford IV antibiotics, or if a patient has difficulties finding someone to administer IV antibiotics, then oral antibiotics are always preferred over no antibiotics. LLMDs can, if necessary, treat a patient with chronic Lyme disease with oral antibiotics for the rest of the patient’s life. Thus, similarities exist between chronic Lyme disease and the Human Immunodeficiency Virus (HIV) that also cannot be eradicated with oral medication.

Up until December 2017, the Centers for Disease Control and Prevention (CDC) in the USA officially endorsed the IDSA treatment guidelines for Lyme disease. What the CDC’s view on treatment for chronic Lyme disease is today is unclear [62]. The CDC still uses the IDSA terminology ‘post-treatment Lyme disease syndrome’ (PTLDS) [63]. It is unlikely that the IDSA’s lack of medical ethics when it comes to Lyme disease [64] was the reason why the CDC stopped officially endorsing IDSA treatment guidelines for Lyme disease, because in 2016 an anonymous whistleblower group
of scientists from within the CDC, calling themselves CDC Scientists Preserving Integrity, Diligence and Ethics in Research (CDC SPIDER), raised concerns about the CDC’s lack of independence and ethics [65], stating, “We are a group of scientists at CDC that are very concerned about the current state of ethics at our agency. It appears that our mission is being influenced and shaped by outside parties and rogue interests”. The CDC therefore most likely stopped endorsing the IDSA treatment guidelines because legal and public pressure was building up. Today people are becoming more aware of chronic Lyme disease. The conflict over what the best ways are to diagnose and treat Lyme disease have gone so far that a Tick-Borne Disease Working Group was created when the 21st-century cures act became law in the USA in December 2016 [66]. The objective of the working group is to review the scientific literature, regarding for example causes, prevention, diagnosis, duration, surveillance and treatment for Lyme disease and associated diseases.

The genus *Borrelia* was named in honor of the French bacteriologist Amédée Borrel in 1907 [67]. The name Lyme comes from the name of the town in Connecticut in the USA where the disease was “first” observed in 1977 [68] despite the fact that the German physician Alfred Buchwald published the first case of atrophy of the skin already in 1883 [69]. In 1902 Herxheimer and Hartmann named such atrophy of the skin Acrodermatitis Chronica Atrophicans (ACA) [70] which is why ACA, which is dermatological disorder mostly associated with chronic Lyme disease in Europe [71], sometimes is called Herxheimer disease. ACA presents itself as a red-bluish discoloration of the extremities [72]. The link between Lyme disease and the bacteria that cause it was first discovered by the medical entomologist Dr. Willy Burgdorfer in 1982 [73]. The bacteria, therefore, took the name after its discoverer *Borrelia* Burgdorferi. It was first in 1929 when the serendipitous discovery of penicillin was made by scientist Alexander Fleming [74] that treatment for Lyme disease became available. Penicillin was originally produced from mold. Naturally occurring antibiotics can also be produced by fermentation, an old technique that can be traced back almost 8000 years [75]. The genes that convey antibiotic resistance to bacteria have been around for at least 30,000 years [76], which means that antibiotic resistance to a large part has not been developed during the last 88 years humankind has known about antibiotics which is a fact that we rarely hear about though.

One of the first people to treat chronic Lyme, or more specifically ACA, with penicillin in 1949 was Nils Thyresson [77] who was a Swedish medical doctor and professor of Dermatology and Venereology. The first human believed to have been infected with Lyme disease was Ötzi the Iceman [78]. This 5300 years old naturally preserved mummy was discovered in 1991 in the Ötztal Alps on the border between Austria and Italy. Spirochete bacteria are believed to be much older this though. Spirochetes similar to *Borrelia* has been found in a 15 million years old tick preserved in amber in the Dominican Republic [79] which means that the spirochete bacteria that causes Lyme disease is much older than humankind. The complete *Borrelia* genome was first sequenced in 1997 [80]. The *Borrelia* bacteria has the most complex genomic architecture among known prokaryotes [81]. *Borrelia* bacteria are also very genetically diversified [82,83]. Different *Borrelia* strains have different antigens [84] which means that different *Borrelia* strains will produce different immune system responses in the form of antibodies in an infected host. A blood test that only test for one specific antibody cannot detect antibodies from different *Borrelia* strains. A person that is infected with one type of *Borrelia* strain will also have different symptoms compared to a person that is infected with a different *Borrelia* strain [85]. For example, arthritis is usually not seen in European Lyme disease patients [86] due to the different *Borrelia* strains that exist in Europe compared to the USA which means that the lack of arthritis in a patient should not be interpreted as the patient not having Lyme disease. Lyme disease diagnosis and treatment are further complicated by the fact that *Borrelia* also has many different co-infections [87] such as Babesia, Bartonella, Ehrlichia, etc. It is believed that a vector such as a tick can spread at least 237 different types of bacteria [88] and many types of viruses [89] which means that many chronic Lyme patients might be infected with many different types of microbes at the same time.
In the Lyme disease documentary Under Our Skin, Dr. Burgdorfer says: “The controversy in the Lyme disease research is a shameful affair and I say this because the whole thing is politically tainted. Money goes to the same people who have for the last thirty years produced the same thing. Nothing.” [90]. The *Borrelia* bacteria has at least three different morphological forms. (1) A spirochete form [91]. (2) A round body form that is also known as cyst form [92]. (3) A biofilm form [93]. The *Borrelia* bacteria evade the immune system by for example changing morphology [94] and by changing its outer surface proteins (Osp) also known as antigens [95–98]. When the bacteria change its antigens all the time, the specific antibodies that are produced by the immune system to try to eradicate the infection becomes useless. The DNA structure that is believed to be responsible for the bacteria’s antigenic variation ability is called G-quadruplex (G4) [99]. Immune system evasion by other diseases such as Amyotrophic Lateral Sclerosis (ALS) [100] and cancer [101] is also believed to be connected to G4. Exciting research is being done that is trying to find medications that block G4 for the *Borrelia* bacteria [102]. If the researchers are successful, it could mean an end to chronic Lyme disease but also to other diseases. If the immune system in combination with some drug could eradicate the *Borrelia* bacteria that would be a superior solution compared to a possible lifetime of oral antibiotics. The bacteria evade being killed by broad-spectrum antibiotics by changing morphology [103]. Note that the bacteria ability to evade broad-spectrum antibiotics should not be interpreted as antibiotic resistant. Metronidazole/Tinidazole forces the bacteria to take a spirochete form [104,105] where other broad-spectrum antibiotics such as Azithromycin can kill it off. When chronic Lyme patients are treated with a macrolide antibiotic such as Azithromycin in combination with Metronidazole/Tinidazole, then the physician can also give Plaquenil which is an anti-malaria medication. Plaquenil raises the PH level in cells so that macrolide antibiotics can work more effectively [106]. Unfortunately, Plaquenil does not work for tetracycline antibiotics [107]. It is, however, important to note that Plaquenil can cause eye problems [108] which means that not all patients can tolerate this medication.

Because different combinations of antibiotics have different effects on the different morphological forms of the *Borrelia* bacteria [109], ILADS treatment guidelines recommend combination therapy with two different types antibiotics for chronic Lyme disease [110] for example metronidazole/tinidazole in combination with azithromycin/doxycycline. The recommended treatment for chronic Lyme disease is therefore different from the recommended treatment for acute Lyme by the IDSA which is, monotherapy with one antibiotic for example doxycycline. Again, similarities exist between HIV and chronic Lyme disease because both are treated with combination therapy. The reason for combination therapy for HIV and chronic Lyme disease is however different. Combination therapy in HIV is motivated by the fact that the virus develops resistance to the medication if you only treat with one antiviral medication. For chronic Lyme disease, this is not the cases. Combination therapy for chronic Lyme disease is motivated by the fact that the *Borrelia* bacteria changes morphology which means that monotherapy is not an effective treatment for Lyme disease.

A common mistake is to describe the symptoms of Lyme disease as “flu-like”. A serious Lyme disease infection should not be reduced to a simple cold. A better screening symptom for Lyme disease is paresthesia. Paresthesia manifests itself as vibrating sensation under the skin. One study estimates that 53% of patients with chronic Lyme disease develop paresthesia [111]. A second study estimates that 70% of patients with chronic Lyme disease develop paresthesia [112]. The good thing is that the paresthesia stops when antibiotics have killed off the infection [113] but also during treatment with antibiotics such as Metronidazole or Tinidazole. There are three main questions that need to be answered to justify treatment for chronic Lyme disease. (1) Do chronic Lyme disease patients without treatment suffer from a low Quality of Life (QOL) than the general population? Four National Institute of Health (NIH) studies have shown that the answer to that question is yes [114]. (2) Are the symptoms of chronic Lyme disease patients reduced because of antibiotic treatment? The answer to that question is yes [115–119]. (3) Can an untreated chronic Lyme disease lead to premature death? The answer to that question is yes according to our previous CDC reference. There exist a few studies that show that the symptoms of chronic Lyme patients do not improve with antibiotics treatment.
There exist at least two problems with these studies. (1) They usually treat chronic Lyme patients with monotherapy which again is not a recommended treatment. (2) They have used flawed statistical methods [120]. Medical doctors should be able to use their professional expertise and in consultation with their patients determine the most appropriate treatment for patients.

A European study [121] estimates the sensitivity of the Lyme disease test to 44%. Such number was found by analyzing the performance of eight different Lyme disease tests. 89 different blood samples were tested for each test manufacturer. The sample also included healthy controls. Another study [122] that conducted a meta-analysis of the American scientific literature (eight scientific articles) regarding the performance of the Lyme disease test estimates the sensitivity of the Lyme disease test to 46% and the specificity to 99%. Because the blood tests that are used today by government health agencies to detect the human body’s production of antibodies against the Borrelia bacteria’s antigens are so insensitive Lyme disease, today must be defined as a clinical diagnosis as correctly advocated by ILADS. If a patient has had a blood test that suggests a Lyme disease infection that is fine, but a positive blood test should not be a requirement for a diagnosis. A person that suspect a Lyme disease infection is today, unfortunately, better of tossing a coin and diagnosing themselves because the blood tests for Lyme disease today will miss most infections. Specific antibodies often cannot be found in patients with Lyme disease because of the bacteria ability to shift its antigens.

Testing cerebrospinal fluid instead of blood will not improve test sensitivity because the bacteria can still shift its antigens in cerebrospinal fluid which means that the probability of detecting specific antibodies in cerebrospinal fluid is not larger than in blood. Moreover, cerebrospinal fluid causes the bacteria to change its morphology from a spirochete form to a cyst from [123]. If you only diagnose Lyme disease in patients that have specific antibodies in their blood or cerebrospinal fluid, you will miss most cases. To better understand Lyme disease testing let’s look at the confusion matrix [124]. All calculations can be found in the macro-enabled excel file “Lyme disease model.xlsx”. We assume in Table 1 that the number of people in the disease and control groups are known and equal. In real life, however, we do not know the number of people in the disease and control group. We can only observe the total number of people that tested positive in both groups.

We can now plot the number of people that tested positive in both groups (z) and the % number of people that tested positive in both groups (zz) when the number of infected people in the disease group (x) and the number of healthy people in control groups (y) are unknown as seen in Figures 1 and 2.

Table 1. Number of people that tested positive in both groups (z) when the number of infected people in the disease group (x) and the number of healthy people in control groups (y) are known and assumed to be equal.

| Assumed values          | Number of infected people in disease group (x) = 100 | Number of healthy people in control group (y) = 100 | Test sensitivity (se) = 0.44 | Test specificity (sp) = 0.99 |
|-------------------------|------------------------------------------------------|-----------------------------------------------------|------------------------------|----------------------------|
| Formulas                | Number of infected people in the disease group that tested positive = true positive (a) = \( x \cdot se \) |
|                         | Number of infected people in the disease group that tested negative = false negative (b) = \( x \cdot a = x \cdot x \cdot se \) |
|                         | Number of healthy people in the control group that tested positive = false positive (c) = \( y \cdot (1-sp) \) |
|                         | Number of healthy people in the control group that tested negative = true negative (d) = \( y \cdot c = y \cdot y \cdot (1-sp) \) |
| Confusion matrix        | Disease group                                      | Control group                                      |                              |
|                         | Number of true positives (a) = 44                  | Number of false positives (c) = 1                 |                              |
|                         | Number of false negatives (b) = 56                 | Number of true negatives (d) = 99                 |                              |
| Statistics              | The relationship between y and x (p) = \( y / x \)  |                                           1 |
|                         | Test sensitivity (se) = \( a / (a + b) \) = 0.44   |                                           0.44 |
|                         | Test specificity (sp) = \( d / (c + d) \) = 0.99   |                                           0.99 |
Table 1. Cont.

| Description                                                                 | Formula                                                                 | Value |
|----------------------------------------------------------------------------|-------------------------------------------------------------------------|-------|
| Number of infected people in the disease group (x)                         | a + b                                                                   | 100   |
| Number of healthy people in the control group (y)                         | c + d                                                                   | 100   |
| Total number of tested people                                              | a + b + c + d                                                           | 200   |
| Number of people that tested positive in both groups (z)                  | (a + c) * x + y * (1-sp) * 100                                          | 45    |
| % of people that tested positive in both groups (zz)                       | (z / (x + y)) * 100                                                    | 22.5  |
| Number of false test results in disease group                             | b                                                                       | 56    |
| % number of false test results in disease group                            | (b / (a + b)) * 100                                                   | 56    |
| Number of correct test results in disease group                            | a                                                                       | 44    |
| % number of correct test results in disease group                          | (a / (a + b)) * 100                                                   | 44    |
| Number of correct test results in control group                            | d                                                                       | 99    |
| % number of correct test results in control group                          | (d / (c + d)) * 100                                                   | 99    |
| Number of false test results in control group                             | c                                                                       | 1     |
| % number of false test results in control group                            | (c / (c + d)) * 100                                                   | 1     |
| Number of false test result in both groups                                 | b + c                                                                   | 57    |
| % number of false test results in both groups                              | (b + c) / (a + b + c + d) * 100                                         | 0.285 |
| % of people that tested positive in disease group                          | (a / (a + b)) * 100                                                   | 44    |

Figure 1. The number of people that tested positive in both groups (z) when the number of infected people in the disease group (x) and the number of healthy people in control groups (y) are unknown.

Figure 2. The % number of people that tested positive in both groups (zz) when the number of infected people in the disease group (x) and the number of healthy people in control groups (y) are unknown.
We can now assume that the number of healthy people in the control group \( y = p \times \) the number of sick people in the disease group \( x \). The relationship between \( y \), \( x \) and \( p \) can be seen in Figure 3.

![Figure 3. The relationship between \( y \), \( x \) and \( p \).](image)

We now can adjust the equation for \( z \) and \( zz \) to include \( p \) as seen in Figures 4 and 5. It can be hard to see in Figure 5 (it is better to look in the excel file directly) but \( zz \) has the same value for all \( x \) values for any given value of \( p \) which means that you can never find a value for \( x \) (and a value for \( y \) thanks to \( y = x \times p \)) only given values for \( se \), \( sp \), \( p \) and \( zz \). I have also written a sub procedure in VBA called \( zzz() \) that also can be found in the excel file “Lyme disease model.xlsx” that can be called by simply pressing a button that calculates \( zz \) given a Lyme disease test sensitivity (\( se \)) equal to 0.44 and a Lyme disease test specificity of (\( sp \)) equal to 0.99.

![Figure 4. The number of people that tested positive in both groups (\( z \)) adjusted for the relationship between \( y \) and \( x \) (\( p \)).](image)
Figure 5. The % number of people that tested positive in both groups (zz) adjusted for the relationship between y and x (p).

We can now solve for the number of infected people in the disease group (x) and the number of healthy people in the control group (y) when they are unknown given a value for z with matrix algebra as seen in Table 2.

Table 2. How to find x and y with matrix algebra.

The equation we need to solve is the following:

\[
\text{Number of people that tested positive in both groups (z)} = \text{the number of true positives (a) + the number of false positives (c)}
\]

where

- Number of people with an infection in disease group: \(x\)
- Number of people without an infection in control group: \(y\)
- Total number of people in the disease and control groups: \(x + y\)
- Test sensitivity (se): 0.44
- Test specificity (sp): 0.99

Number of true positives (a) = \(x \times \text{se}\)

\(\begin{align*}
\text{Number of false positives (c) = } & y \times (1 - \text{sp}) \\
\text{Number of people that tested positive in both groups (z) = } & 100,000
\end{align*}\)

We can solve such equation in excel by using matrix algebra. The system of linear equations we should solve is

\[
A \times B = C
\]

where

\[
\begin{align*}
A &= \begin{bmatrix} 0.44 & -2.5 \\ -2.5 & 1 \end{bmatrix} \\
B &= \begin{bmatrix} x \\ y \end{bmatrix} \\
C &= \begin{bmatrix} 0 \\ 100,000 \end{bmatrix}
\end{align*}
\]

\[
A^{-1} = \begin{bmatrix} 2.150538 & -0.021505 \\ 5.376344 & 0.946237 \end{bmatrix}
\]

\[
B = A^{-1} \times C
\]

\[
\begin{align*}
x &= 215,054 \\
y &= 537,634
\end{align*}
\]
Table 2. Cont.

| The relationship between y and x (p) = y / x | 2.5 |
| Number of people that test positive in the disease group = true positive (a) = x * se | 94,624 |
| Number of people that test positive in the control group = false positive (c) = y * (1-sp) | 5,376 |
| Number of people that tested positive in both groups (z) = a + c = x * se + y * (1-sp) | 100,000 |
| % of people that tested positive in both groups (zz) = (z / (x + y)) * 100 | 13.29 |

I have written a user defined function (udf) in VBA that does the above calculations automatically.

Lyme1(se ; sp ; p ; z ; output) where output is either “x”, “y” or “zz”

| Lyme1(0.44;0.99;2.5;100000;”x”) | x = 215,054 |
| Lyme1(0.44;0.99;2.5;100000;”y”) | y = 537,634 |
| Lyme1(0.44;0.99;2.5;100000;”zz”) | zz = 13.29 |

We can algebraically manipulate the previous equations further as seen in Table 3.

Table 3. Algebraic manipulation of previous equations.

| We know that | We know that |
|---|---|
| z = x * se + y * (1-sp) | z = x * se + y * (1-sp) |
| y = p * x | x = y / p |
| which means that | which means that |
| z = x * se + p * x * (1 - sp) | z = (y / p) * se + y * (1 - sp) |
| we solve for x | we solve for y |
| x = z / (-sp * p + se + p) | y = p * z / (-sp * p + se + p) |

For the previous example with z = 100,000 and p = 2.5 we get

| x = 215,054 |
| y = 537,634 |

I have again written a udf in VBA.

Lyme2(se ; sp ; p ; z ; output) where output is either “x”, “y” or “zz”

| Lyme2(0.44;0.99;2.5;100000;”x”) | x = 215,054 |
| Lyme2(0.44;0.99;2.5;100000;”y”) | y = 537,634 |
| Lyme2(0.44;0.99;2.5;100000;”zz”) | zz = 13.29 |

we know that

| zz = (z / (x + y)) * 100 | zz = (z / (x + y)) * 100 |
| y = p * x | x = y / p |
| which means that | which means that |
| zz = (z / (x + p * x)) * 100 | zz = (z / ((y / p) + y)) * 100 |
| we solve for x | we solve for y |
| x = 100 * z / (p * zz + zz) | y = 100 * p * z / (p * zz + zz) |
Table 3. Cont.

For the previous example with $z = 100,000$, $zz = 13.29$ and $p = 2.5$ we get:

$$x = 215,054$$
$$y = 537,634$$

I have again written a udf in VBA:

Lyme3(se; sp; p; z; zz; output) where output is either "x", "y"

Lyme3(0.44;0.99;2.5;100000;13.29;"x")

$$x = 215,054$$

Lyme3(0.44;0.99;2.5;100000;13.29;"y")

$$y = 537,634$$

We can do some further modeling and plotting as seen in Table 4, Figures 6 and 7.

Table 4. The number of infected people in the disease group ($x$) given total number of people ($T$) and the relationship between $x$ and $y = p$.

| Number of infected people in disease group | $x$ |
| Number of healthy people in control group | $y$ |
| Total number of people ($T$) | $x + y$ |

- $T = x + y \rightarrow y = T - x$
- $y = p \times x$

This means that:

- $T - x = p \times x$
- $T - y = y / p$

We can solve for $x$:

- $x = T / (p + 1)$

We can solve for $y$:

- $y = p \times T / (p + 1)$

Figure 6. The number of infected people in the disease group ($x$) given total number of people ($T$) and the relationship ($p$) between $x$ and $y$. 
Figure 7. The number of healthy people in the control group (y) given the total number of people (T) and the relationship (p) between x and y.

2. Methods

The first limitation of this study is that we are using the incidence rate of Lyme disease from the USA on the population of Europe to get an estimate on how frequent Lyme disease is in Europe. Using the incidence rate from the USA on the population of Europe is not optimal but unfortunately, our only choice since the ECDC do not have any data on Lyme disease. Lyme disease is not spread exclusively by vectors such as ticks. One study [125] explains that Lyme disease is also a Sexually Transmitted Disease (STD). Since Lyme disease can spread sexually, the growth rate of the disease will be similar in the USA and Europe which means that it's not unrealistic to use the disease frequency from the USA on Europe's population. The second limitation of this study is that scientific literature that exists regarding Lyme disease as an STD is very limited. In a perfect world more, research should be done regarding Lyme disease as an STD but please keep in mind that Borrelia's close cousin the spirochete bacterium *Treponema pallidum* that causes syphilis is well known to spread sexually. Hence, the fact that the *Borrelia* bacteria can also or could also spread through sexual contact should not come as a surprise. Unfortunately, accurately modeling the spread of Lyme disease without sexual transmission is close to impossible because there are so many difficult questions that need to be answered regarding tick ecology [126] for example in which geographic areas are ticks most frequent?, how many different types of ticks exists?, which types of ticks spread disease?, what types of microorganisms do ticks contain?, how often do ticks reproduce?, when they reproduce how many eggs do they produce on average?, how many of these eggs survive?, what is the probability that a tick bites a human?, what is the probability that a tick bites an animal?, what is the probability of a transmission of *Borrelia* or any other microorganism after a tick bite?, what animals do ticks use for transportation deer, bird or rodent, how many of these animals that transport ticks are there in a given geographic area?, how is a tick population affected by cold temperature like a winter? As I said, the epidemiological modeling becomes close to impossible especially for an economist maybe not for a biologist.

All infectious diseases (total number of infections in a population) that are transmitted from person to person and where treatment is inadequate has by nature exponential growth over time. A pandemic initially starts with 1 infected person (total number of infections at t1 = 1) that person then goes and infect 1 more person (total number of infections at t2 = 2), these 2 infected people then go and infect 2 more people (total number of infections at t3 = 4), these 4 people then go and infect 4 more
people (total number of infections at \( t = 4 = 8 \)) etc. The % change at \( t = 2 = (2 - 1)/1 \times 100 = 100 \), the % change at \( t = 3 = ((4 - 2)/2 \times 100 = 100 \), the % change at \( t = 4 = ((8 - 4)/4 \times 100 = 100 \). Since the percentage change is constant over time, exponential growth models are also called constant growth rate models.

For a linear growth model, the percentage change is decreasing over time. The relationship between the transmission rate of an infection that is spread through sexual contact and the annual growth rate of infection can be seen in Table 5. We can see that given that each infected person has sex with one healthy person each year then the transmission rate is equal to the annual percentage growth rate of infection for an infectious disease that is spread sexually. The third limitation of this study is that we assume that the annual growth rate for Lyme disease is 2%. To my knowledge, there does not exist any scientific literature regarding the annual growth rate of Lyme disease. The 2% number is therefore not based on empirical observation. However, we can compare such number to other infectious diseases that are also spread through sexual contact such as HIV. According to the CDC, HIV in 1977 had a transmission rate of 100% and in 2006 the transmission rate of HIV was 5% [127]. It becomes obvious that 2% is a very conservative number. The annual growth rate of Lyme disease is most likely higher than 2% because unfortunately many Lyme disease patients and especially chronic Lyme patients are struggling to find antibiotic treatment but since I lack a scientific reference for my claim and because I do not want to inflate the numbers I have chosen to report 2%. The fourth limitation of this study is that the scientific literature regarding how successful IV treatment is at curing chronic Lyme disease is very limited. However, the exact cure rate is not that important because we have known for a long time that antibiotics kill bacteria. We can simply assume that the cure rate is unknown and compare the outcome of five different assumed cure rates and scenarios: 0%, 25%, 50%, 75% and 100%.

### Table 5. The transmission rate for an STD and its relationship to the annual growth rate of infection.

| \( g \) | \( n \) | \( t \) | \( tni1 \) | \( tni2 \) | \( \% \text{change} \) |
|-------|-------|-------|-------|-------|-------|
| \( 1 \) | 1 | 1 | 1 | 1 | 1 |
| \( 2 \) | 2 | 100 | 2 | 100 | 3 |
| \( 3 \) | 4 | 100 | 3 | 100 | 9 |
| \( 4 \) | 8 | 100 | 4 | 100 | 27 |
| \( 5 \) | 16 | 100 | 5 | 100 | 81 |
| \( 6 \) | 32 | 100 | 6 | 100 | 243 |
| \( 7 \) | 64 | 100 | 7 | 100 | 729 |

\( g = \frac{\% \text{change}}{100} \)

\( n = \frac{\% \text{change}}{100} \)

\( t = \frac{\% \text{change}}{100} \)

\( \text{tni1(y)} = \text{tni1(y-1)}^*(1+g/100) \)

\( \text{tni2(y)} = \text{tni2(y-1)}^*(1+t/100) \times n \)

We can see that given that each infected person has sex with one healthy person each year then the transmission rate is equal to the annual percentage growth rate of infection.
3. Results

According to one scientific study [128], that can be found on the CDC website and published in Clinical Infectious Diseases which is a “scientific” journal issued by the IDSA. 2.4 million individual Lyme disease blood tests were done in the USA 2008. That number comes from a survey that was sent to seven of the largest commercial laboratories in the USA. The most interesting part of such publication an equation was not included in the article itself but was present in the supplementary data [129]. The equation that the entire paper is based on is:

\[
\text{Observed } \% \text{ positive } = \% \text{ True Infection } \times \text{ Sensitivity} + (1 - \% \text{ True Infection}) \times (1 - \text{Specificity})
\]

If we multiply the above equation by the number of individual Lyme disease blood tests that were performed in 2008 in the USA, then we get the observed number of Lyme disease infections in the USA in 2008 by the CDC.

\[
\text{Observed positive } = (\% \text{ True Infection } \times \text{ Sensitivity} + (1 - \% \text{ True Infection}) \times (1 - \text{Specificity})) \times \text{ Number of Lyme disease tests}
\]

We solve the first equation for \% True infection.

\[
\% \text{ True Infection } = \frac{(\text{Observed } \% \text{ Positive } + \text{Specificity } - 1)}{(\text{Specificity } + \text{Sensitivity } - 1)}
\]

If we multiply the above equation by the number of individual Lyme disease blood tests that were performed in 2008 in the USA, then we get CDC’s predicted number of Lyme disease infections in the USA in 2008.

\[
\text{True Infection } = \left(\frac{(\text{Observed } \% \text{ Positive } + \text{Specificity } - 1)}{(\text{Specificity } + \text{Sensitivity } - 1)}\right) \times \text{ Number of Lyme disease tests}
\]

The CDC study estimated three different scenarios: high, low and average with an assumed test sensitivity of 66.9% and assumed specificity of 96.1%. Unfortunately, the only value for observed % positive that the CDC provides was 11.89%, which was the average scenario. The other two are not reported in the paper and need to be calculated. Why is the observed % positive variable so important for our calculations later? The observed % positive value has not been adjusted for the sensitivity and specificity of the Lyme disease blood test. Therefore, we can use our values for sensitivity and specificity. The observed % positive values do also not include any assumptions about the relationship between the number of sick people in the disease group (x) and the number of healthy people in the control group (y) that we previously defined as p. The CDC calculations can be found in Table 6.

| CDC equations                                                                                                                                   |
|-----------------------------------------------------------------------------------------------------------------------------------------------|
| Observed % Positive = % True Infection × Sensitivity Lyme test + (1 - % True Infection) × (1 – Specificity Lyme test)                       |
| Observed positive = (% True Infection × Sensitivity + (1 - % True Infection) × (1 – Specificity)) × Number of performed individual Lyme disease tests |
| % True Infection = (Observed % Positive + Specificity – 1) / (Specificity + Sensitivity – 1)                                                   |
| True Infection = ((Observed % Positive + Specificity – 1) / (Specificity + Sensitivity – 1)) × Number of performed individual Lyme disease tests |
| The performed number of individual Lyme disease test in 2008 in the USA (n) 2 400 000                                                        |
| Lyme disease test sensitivity assumed by the CDC (CDC se) 0.669                                                                            |
| Lyme disease test specificity assumed by the CDC (CDC sp) 0.961                                                                            |

Table 6. Comparison 1 of the CDC model and my Lyme disease model.
Table 6. Cont.

| CDC scenario | Low       | Average    | High       |
|--------------|-----------|------------|------------|
| % true infection also known as predicted % positive estimated by the CDC (CDC xx) | 0.1       | 0.1189     | 0.185      |
| Observed % positive in the control and disease group according to or implied by the CDC (CDC zz) | 0.102     | 0.1189     | 0.15555    |
| The number of people that tested positive in the control and disease group in the USA in 2008 according to the CDC (observed positive or CDC x) | 244 800   | 285 360    | 373 320    |
| The number of infected people in the disease group estimated by the CDC (predicted positive also known as true infection or CDC x) | 240 000   | 285 360    | 444 000    |
| CDC z = CDC x * CDC se + CDC y * (1 - CDC sp) | 244 800 = 240 000 * 0.669 + y * (1-0.961) | 285 360 = 285 360 * 0.669 + y * (1-0.961) | 373 320 = 444 000 * 0.669 + y * (1-0.961) |
| The number of healthy people in the control group according the CDC (CDC y) | 2 160 000 | 2 421 901 | 1 965 000 |
| The relationship between y and x = CDC p = CDC y / CDC x | 9.0000    | 8.4872     | 4.4257     |
| Number of healthy people in the control group (y) : number of infected people in the disease group (x) | 9 : 1     | 8.4872 : 1 | 4.4257 : 1 |

My model

We assume p = 1, se = 0.44, sp = 0.99 and z = CDC z in Lyme2(se ; sp ; p ; z ; Output = q) where q is either "x", "y" or "zz"?

| x  | y  | zz |
|----|----|----|
| 544 000 | 634 133 | 829 600 |
| 22.50 | 22.50 | 22.50 |

We assume p = 0.5, se = 0.44, sp = 0.99 and z = CDC z in Lyme2(se ; sp ; p ; z ; Output = q) where q is either "x", "y" or "zz"?

| x  | y  | zz |
|----|----|----|
| 550 112 | 641 258 | 838 921 |
| 275 056 | 320 629 | 419 461 |
| 29.67 | 29.67 | 29.67 |

We can see that the CDC has unrealistically and secretly assumed without motivation or explanation that the size of the control group with healthy people is 9 (low scenario), 8.5 (average scenario) and 4.4 (high scenario) times the size of the disease group with infected people (p = 9, p = 8.5 and p = 4.4). These assumed p-values cannot be empirically observed because it is impossible to know the value of p by simply looking at the total number of people that test positive in the control and disease group (z). It would be more realistic to assume that the size of the control group with healthy people is the same size as the disease group with infected people (p = 1). Because blood testing today is so primitive (you can more or less only test for one pathogen at the time) and because the blood tests for Lyme disease are so insensitive most people today will only test to get a confirmation that they are infected. Hence, many people will only test if they are infected. It is therefore unrealistic to expect that a lot of people without an infection will demand to get tested just for “fun” because there is also a personal financial cost involved. The CDC also claims in their paper that the value of the variable observed % positive was the same as the value for the variable predicted % positive for the average scenario which is mathematically impossible as seen in Figure 8 using the same assumptions regarding Lyme disease test sensitivity and specificity as the CDC. Such claim by the CDC, therefore, does not make any sense.
Figure 8. The value of observed % positive cannot be equal to predicted % positive for the average scenario in the CDC model.

We can now evaluate whether the CDC equation is correct and compare the CDC model with my model as seen in Table 7.

Table 7. Comparison of the CDC model and my Lyme disease model.

| CDC scenario | Low       | Average   | High      |
|--------------|-----------|-----------|-----------|
| The number of infected people in the disease group estimated by the CDC (predicted positive also known as true infection or CDC x) | 240,000   | 285,360   | 444,000   |
| CDC assumptions | CDC z = 244,800 | CDC z = 285,360 | CDC z = 373,320 |
| Assumptions | CDC z = 244,800 | CDC z = 285,360 | CDC z = 373,320 |
| Lyme2(se ; sp ; p ; z ; output) = Lyme2(0.669 ; 0.961 ; 1 ; CDC z ; 'x') | 345,763   | 403,051   | 527,288   |
| CDC assumptions | CDC p = 9 and CDC z = 244,800 | CDC p = 8.4872 and CDC z = 285,360 | CDC p = 4.4257 and CDC z = 373,320 |
| (A) Lyme2(0.669 ; 0.961 ; CDC p ; CDC z ; 'x') | 240,000   | 285,360   | 443,582   |
| CDC x - A | 0         | 0         | 418       |
| (B) Lyme2(0.44 ; 0.99 ; 1 ; CDC z ; 'x') | 533,333   | 634,133   | 829,600   |
| B - A | 293,333   | 348,773   | 386,018   |
| % difference between A and B = (B - A)/B | 55        | 55        | 47        |

Given the CDC unrealistic assumptions regarding se, sp and p, it appears that the equation is more or less correct. There is, however, an unexplained difference of +418 infections for the high scenario. We can also see that if the CDC had assumed that p = 1 in combination with the CDC assumptions se = 0.669, sp = 0.961, then the number of Lyme disease infections in the USA for 2008 for the high scenario would have been 527,288 instead of 444,000. We can also see that the difference between the CDC model with CDC assumptions of se = 0.669, sp = 0.961, p(low) = 9, p(average) = 8.5 and p(high) = 4.4 and my model with my assumptions of se = 0.44, sp = 0.99 and p = 1 is lowest for the CDC high scenario hence I will report that in 2008 in the USA the estimated number of Lyme disease infections is 829,600. The data and calculations that I will present now can be found in the excel file “Lyme disease calculations.xlsx.” Given 829,600 Lyme disease infections in 2008 in the USA and an assumed annual infection growth rate of 2% then in 2018 in the USA approximately 1 million people will get infected with Lyme disease. The total number of Lyme disease infections that we can expect in the USA from 2008 to 2050 is 55.7 million. Given an annual population growth rate in the USA of 1% and given that everyone that is infected from 2008 to 2050 is still alive than 12% of the population in the USA will have been infected with Lyme disease in 2050. The plots are presented in Figures 9–12.
Figure 9. Number of Lyme disease infections in the USA between the years 2008 to 2050.

Figure 10. Annual incidence rate of Lyme disease in the USA between the years 2008 to 2050.

Figure 11. Total number of Lyme disease infections in the USA between the years 2008 to 2050.
Figure 12. % of the population in the USA that have been infected with Lyme disease between the years 2008 to 2050.

Since the ECDC does not have any statistics on how frequent Lyme disease is in Europe, it is difficult to determine an exact figure for the number of Lyme disease cases in Europe in 2008. However, we can use the incidence rate (frequency) from the USA for 2008 on Europe’s population for 2008 to get an estimate. The estimated number of Lyme disease infections in Europe in 2008 is 2,008,505. Given 2,008,505 Lyme disease infections in 2008 in Europe and an assumed annual infection growth rate of 2% then in 2018 in Europe approximately 2.4 million people will get infected with Lyme disease. The total number of Lyme disease infections that we can expect in Europe from 2008 to 2050 is 134.9 million. Given an annual population growth rate in Europe of 0.2% and given that everyone that is infected from 2008 to 2050 is still alive than 17% of the population of Europe will have been infected with Lyme disease in 2050. The plots are presented in Figures 13–16.

Figure 13. Number of Lyme disease infections in Europe between the years 2008 to 2050.
According to our previous scientific reference, 63% of people that are infected with *Borrelia* develop chronic Lyme disease. The estimated number of people suffering from chronic Lyme disease in 2050 in the USA and Europe depend on the assumption we make regarding the estimated cure rate for chronic Lyme disease treatment and on the assumption that everyone that develop chronic Lyme disease from 2008 to 2050 is still alive. The last assumption is very uncertain because of how difficult it is today for chronic Lyme patients to get treatment. We can see in Table 8 and Figure 17 that the number of people that will suffer from chronic Lyme disease in 2050 in the USA with a 0% cure rate is approximately 35 million (8% of the USA population in 2050), with a 25% cure rate 26 million (6% of the USA population in 2050), with a 50% cure rate 18 million (4% of the USA population in 2050), with a 75% cure rate 9 million (2% of the USA population in 2050) and with a 100% cure rate zero (0% of the USA population in 2050).
Figure 16. % of the population in Europe that have been infected with Lyme disease between the years 2008 to 2050.

Table 8. Number of people with chronic Lyme disease in 2050 in the USA.

| Assumed cure rate | Number of people with chronic Lyme disease in 2050 in the USA | % of USA’s population with chronic Lyme disease in 2050 |
|-------------------|-------------------------------------------------------------|------------------------------------------------------|
| 0% cure rate      | 35 100 762                                                  | 8                                                   |
| 25% cure rate     | 26 325 571                                                  | 6                                                   |
| 50% cure rate     | 17 550 381                                                  | 4                                                   |
| 75% cure rate     | 8 775 190                                                   | 2                                                   |
| 100% cure rate    | 0                                                           | 0                                                   |
We can see in Table 9 and Figure 18 that the number of people that will suffer from chronic Lyme disease in 2050 in Europe with a 0% cure rate is approximately 85 million (11% of Europe’s population in 2050), with a 25% cure rate 64 million (8% of Europe’s population in 2050), with a 50% cure rate 42 million (5% of Europe’s population in 2050), with a 75% cure rate 21 million (3% of the Europe’s population in 2050) and with a 100% cure rate zero (0% of Europe’s population in 2050).

Table 9. Number of people with chronic Lyme disease in 2050 in Europe

| Assumed cure rate | Number of people with chronic Lyme disease in 2050 in Europe | % of Europe’s population with chronic Lyme disease in 2050 |
|-------------------|-------------------------------------------------------------|----------------------------------------------------------|
| 0% cure rate      | 84 980 791                                                  | 11                                                       |
| 25% cure rate     | 63 735 593                                                  | 8                                                        |
| 50% cure rate     | 42 490 396                                                  | 5                                                        |
| 75% cure rate     | 21 245 198                                                  | 3                                                        |
| 100% cure rate    | 0                                                           | 0                                                        |

If 2,008,505 people were infected with Lyme disease in Europe in 2008 and go undetected by the non-exist statistics what types of illnesses can Lyme disease patients be misdiagnosed with? It is therefore not unrealistic to assume that some Lyme disease patients are misdiagnosed with illnesses such as for example fibromyalgia, chronic fatigue syndrome (CFS), heart arrhythmia, multiple sclerosis (MS), restless legs syndrome (RLS), amyotrophic lateral sclerosis (ALS), Parkinson disease, systemic lupus erythematosus (SLE), Gulf War Syndrome (GWS) or Alzheimer’s disease. The commonality between these diseases is that there is currently no known cure. Before we can start to estimate costs for Lyme disease, we need to discuss the tax on disability benefits. The tax rate on disabilities benefits is approximately 50% of the tax rate on income from work [130,131]. However, the tax
on disability benefits is irrelevant to the government because assuming the government keeps the amount of disability benefit after tax fixed an increased tax rate on disability benefits will not increase the government’s financial balance (government revenues + government costs) as seen in Table 10. If a government raises the tax rate on a person’s disability benefit, then the increased tax revenues from such tax raise are completely offset by the larger government cost for the disability benefit.

![Figure 18. Number of people with chronic Lyme disease in 2050 in Europe.](image)

**Table 10.** A government’s financial balance (government revenues + government costs) with a disability benefit tax increase.

| A government’s financial balance after tax for a disabled person not working without the lost tax revenue because a chronic Lyme patient is not working |  |
| --- | --- |
| Annual disability benefits after tax (ADAT) | 13 000 |
| % tax rate on income from work | 0.40 |
| Tax rate in % on disability benefits | 0.20 |
| Annual disability benefits before tax T1 = ADAT / (1 - T1) | 16 250 |
| Difference | 21 667 |
| Government revenues | 3 250 |
| Tax revenues from disability benefits (+) | 8 667 |
| Government costs for a disabled person | -16 250 |
| Disability benefits (-) | -21 667 |
| A government’s financial balance = tax revenue + government cost | -13 000 |
| | -13 000 |
| Difference | 0 |

| A government’s financial balance for an enabled person working vs a disabled person not working before tax |  |
| --- | --- |
| Annual disability benefits before tax (ADAT) | 13 000 |
| Annual income from work before tax | 35 000 |
| % tax rate on income from work | 0.4 |
| A government’s financial balance = tax revenue + government cost | 14 000 |
| An enabled person that is working | A disabled person that is not working | Difference |
| Government revenues | 14 000 | 0 | -14 000 |
| Tax revenues from income from work (+) | 14 000 | 0 | -14 000 |
| Government costs for a disabled person (-) | 0 | -13 000 | -13 000 |
| Disability benefits (-) | 0 | -14 000 | -14 000 |
| Lost tax revenues because a disabled person is not working (-) | 0 | -14 000 | -14 000 |
| Total costs | 0 | -27 000 | -27 000 |
| A government’s financial balance = tax revenue + government cost | 14 000 | -27 000 | -41 000 |
We can now estimate the costs for Lyme disease treatment in the USA and Europe for the year 2018. We have previously estimated that the number of infections in the USA for 2018 is approximately 1 million. In Europe, that number is approximately 2.4 million. Let’s assume that in the USA for 2018 the annual cost for oral antibiotics is 1400 USD and the annual cost for IV antibiotics is 15,000 USD. In Europe, the estimated annual cost for oral antibiotics is 1200 EUR and the annual cost for IV antibiotics is 13,000 EUR. Currently, the treatment cost for IV antibiotics for chronic Lyme disease are paid by the individual and not by the government or insurance companies. Note that the cost of IV treatment for chronic Lyme disease can be different in the USA and Europe because in the USA a peripherally inserted central catheter (PICC) is preferred while in Europe an IV drip is preferred. The cost for IV treatment in Europe could easily be twice as large because patients must travel a long way and stay at a hotel close to a specialized chronic Lyme disease clinic to get daily IV treatments with a butterfly needle. Sometimes people are even forced to sell their house to afford treatment. We assume that acute Lyme disease is treated with oral antibiotics for one month and that chronic Lyme disease is treated with IV antibiotics for six months or one year. Chronic Lyme disease can also be treated with oral antibiotics but then the patient runs the risk of having to take oral antibiotics for the rest of his/her life. I have therefore not calculated the cost of treating chronic Lyme with oral antibiotics since its more realistic to assume that all chronic Lyme patients want to treat with IV antibiotics because then they will at least have a chance of getting cured of their “chronic” infection. We can see in Tables 11 and 12 that the estimated treatment cost for Lyme disease for 2018 for the USA is somewhere between 4.8 billion USD and 9.6 billion USD and for Europe somewhere between 10.1 billion EUR and 20.1 billion EUR depending on the assumptions we make regarding the length of treatment with IV antibiotics (six months vs. one year). We can also see that the cost of treating acute Lyme disease with oral antibiotics for one month only represents 0.6% of the average treatment cost for IV antibiotics. The cost of treating acute Lyme disease with oral antibiotics is so small that it barely has an impact on total treatment costs.

Table 11. The estimated total cost for the treatment of acute and chronic Lyme disease for the USA for 2018.

| Treatment costs for Lyme disease for the USA for 2018 | Values       |
|------------------------------------------------------|--------------|
| Number of infections 2018 in the USA                 | 1 011 278    |
| People infected with Lyme that develop chronic Lyme | 0.63 %       |
| People infected with Lyme that develop acute infection| 0.37 %       |
| The assumed annual cost of oral antibiotics is       | 1 400 USD    |
| The assumed annual cost of IV antibiotics is         | 15 000 USD   |
| The cost of treating acute Lyme disease with oral antibiotics for one month for the USA for 2018 (Oral USA 2018) | 43 653 490 44 million USD |
| The cost of treating chronic Lyme disease with IV antibiotics for 0.5 years for the USA for 2018 (IV0.5 USA 2018) | 4 778 287 467 4.8 billion USD |
| The cost of treating chronic Lyme disease with IV antibiotics for 1 year for the USA for 2018 (IV1 USA 2018) | 9 556 574 934 9.6 billion USD |
| Total cost of treating acute with oral antibiotics for one month and chronic with IV antibiotics 0.5 years for the USA for 2018 | 4 821 940 958 4.8 billion USD |
| Total cost of treating acute with oral antibiotics for one month and chronic with IV antibiotics 1 year for the USA for 2018 | 9 600 228 425 9.6 billion USD |
| Oral USA 2018 / ((IV0.5 USA 2018 + IV1 USA 2018)/2) | 0.6 %        |
Table 12. The estimated total cost for the treatment of acute and chronic Lyme disease for Europe for 2018.

| Treatment costs for Lyme disease for Europe for 2018 | Values                  |
|----------------------------------------------------|-------------------------|
| Number of infections 2018 in Europe                | 2,448,357               |
| People infected with Lyme that develop chronic Lyme | 0.63%                   |
| People infected with Lyme that develop an acute infection | 0.37%                   |
| The assumed annual cost of oral antibiotics is      | 1,200 EUR               |
| The assumed annual cost of IV antibiotics is        | 13,000 EUR              |

The cost of treating acute Lyme disease with oral antibiotics for one month for Europe in 2018 (Oral Europe 2018) 90,589,198 91 million EUR
The cost of treating chronic Lyme disease with IV antibiotics for 0.5 years for Europe in 2018 (IV0.5 Europe 2018) 10,026,020,721 10.0 billion EUR
The cost of treating chronic Lyme disease with IV antibiotics for 1 year for Europe in 2018 (IV1 Europe 2018) 20,052,041,441 20.1 billion EUR
Total cost of treating acute with oral antibiotics for one month and chronic with IV antibiotics 0.5 years for Europe in 2018 10,116,609,919 10.1 billion EUR
Total cost of treating acute with oral antibiotics for one month and chronic with IV antibiotics 1 year for Europe in 2018 20,142,630,639 20.1 billion EUR
Oral Europe 2018 / ((IV0.5 Europe 2018 + IV1 Europe 2018)/2) 0.6 %

We have so far only looked at treatment cost. Many people are forced to leave their jobs due to the infection. So, the cost of lost personal income is also high. According to one study, approximately 42% of chronic Lyme disease patients cannot work [132]. I believe that the number is even higher but 42% is reasonable enough. The financial cost for both the individual (lost earnings from not working due to chronic Lyme disease) and for the government (lost tax revenues because chronic Lyme patients are not working and disability benefits for chronic Lyme disease patients) are therefore significant. We can now calculate the government cost for chronic Lyme disease for the USA and Europe for 2018 if governments do not finance treatment. According to the Organisation for Economic Co-Operation and Development (OECD) the average annual wage rate before tax for 2016 for the USA was 60,154 USD and for Germany 38,302 EUR [133]. The average annual wage rate in Germany represents the average annual wage rate for Europe. The global accounting firm KPMG states that the income tax rate for 2016 for the USA is 39.6% and for Germany 45% [134]. A website called disabilitysecrets.com [135] claims that the average monthly Social Security Disability Insurance (SSDI) payment after tax for the USA for 2017 is 1,171 USD which means that the average annual disability benefit after tax for the USA for 2017 is 14,052 USD. This amount is on par with the lowest monthly disability payment after taxed paid in Sweden which is 11,000 Swedish kroner [136] which means that the minimum annual disability benefit after tax for Sweden is 132,000 Swedish kroner which as of October 2017 is approximately 13,778 EUR. We will use the monthly Swedish disability payment to represent the disability payment in Europe.

We can see in Tables 13 and 14 that the estimated lost personal income for chronic Lyme patients that are not working for 2018 for the USA is 16.1 billion USD and in Europe 24.8 billion EUR. If the governments in the USA and Europe do not finance IV treatment with antibiotics for chronic Lyme disease, then the estimated government costs for chronic Lyme disease (lost tax revenues because some chronic Lyme patients are not working plus disability benefits for chronic Lyme patients that not working) for 2018 for the USA is 10.1 billion USD and for Europe 20.1 billion EUR. We can also see that the government cost for chronic Lyme disease for 2018 (which does not include treatment costs) in the USA is 2.1 times larger than the cost for 6 months of IV treatment with antibiotics in the USA for 2018 and 1.1 times larger than the cost for 1 year of IV treatment with antibiotics in the USA for 2018. In Europe, the government cost for chronic Lyme disease for 2018 is two times larger than the cost for six months of IV treatment in Europe for 2018 and the government cost is on par with the cost for one year of IV treatment in Europe for 2018.
Table 13. Government cost for chronic Lyme disease for the USA for 2018 if governments do not finance IV treatment with antibiotics for chronic Lyme disease.

| Government costs for chronic Lyme disease for the USA for 2018 if the government does not finance IV treatment for chronic Lyme disease | Values |
|---|---|
| Number of Lyme disease infections 2018 in the USA | 1 011 278 |
| People infected with Borrelia that develop chronic Lyme | 637 105 |
| Number of chronic infections in the USA for 2018 | 0.63 % |
| % of chronic Lyme patients that does not work | 0.42 % |
| % of chronic Lyme patients that does work | 0.58 % |
| Average annual income from work before tax for the USA for 2016 | 60 154 USD |
| Income tax rate for the USA for 2016 | 0.396 % |
| Average annual disability benefits after tax for the USA for 2017 | 14 052 USD |

Lost personal income for chronic Lyme patients not working in the USA for 2018: 16 096 253 841 billion USD

Lost tax revenues because chronic Lyme patients are not working in the USA in 2018 (LTR USA 2018): 6 374 116 521 billion USD

Disability benefits for chronic Lyme patients for the USA for 2018 (DB USA 2018): 3 760 091 747 billion USD

Government cost for chronic Lyme disease for the USA in 2018 (GCL USA 2018) = LTR USA 2018 + DB USA 2018: 10 134 208 268 billion USD

GCL USA 2018 / IV0.5 USA 2018: 2.1

GCL USA 2018 / IV1 USA 2018: 1.1

Table 14. Government cost for chronic Lyme disease for Europe for 2018 if governments do not finance IV treatment with antibiotics for chronic Lyme disease.

| Government costs for chronic Lyme disease for Europe for 2018 if the governments do not finance IV treatment for chronic Lyme disease | Values |
|---|---|
| Number of infections 2018 in Europe | 2 448 337 |
| People infected with Borrelia that develop chronic Lyme | 1 542 465 |
| Number of chronic infections in Europe for 2018 | 0.63 % |
| People that cannot work due to chronic Lyme | 0.42 % |
| Average annual income from work before tax for Germany for 2016 | 38 302 EUR |
| Income tax rate for Germany for 2016 | 0.45 % |
| The minimum annual disability benefits after tax in Sweden for 2016 is 11 000 Swedish kroner | 13 778 EUR |

Lost personal income for chronic Lyme patients not working in Europe in 2018: 24 813 383 257 billion EUR

Lost tax revenues because chronic Lyme patients are not working in Europe for 2018 (LTR Europe 2018): 11 166 022 466 billion EUR

Disability benefits for chronic Lyme patients in Europe for 2018 (DB Europe 2018): 8 925 976 833 billion EUR

Government cost for chronic Lyme disease for Europe for 2018 (GCL Europe 2018) = LTR Europe 2018 + DB Europe 2018: 20 091 999 298 billion EUR

GCL Europe 2018 / IV0.5 Europe 2018: 2.0

GCL Europe 2018 / IV1 Europe 2018: 1.0

Today’s and future government revenues and costs for chronic Lyme disease are however much more important to look at than historical annual cost. We will again treat the cure rate for IV antibiotics for chronic Lyme disease as an unknown. We will calculate variables for five different scenarios: A cure rate of 0%, a cure rate 25%, a cure rate of 50%, a cure rate of 75% and a cure rate of 100%. The number of chronic Lyme patients that are cured with treatment/not cured with treatment and are working/not working can be seen in Table 15.
Table 15. The work/not work/cured/not cured matrix.

| % cure rate (cr) | Cured from treatment | Not cured from treatment | Sum |
|------------------|----------------------|--------------------------|-----|
| Number of people | C = nn * (cr/100)    | NC = nn - C              | C + NC |
| Number of people that are working | C * 1 | NC * (w / 100) | C * 1 + NC * (w / 100) |
| Number of people that are not working | C * 0 | NC * (nw / 100) + | C * 0 + NC * (nw / 100) + |
| Sum | C+C+0 | NC*(nw/100) | C*0+NC*(nw/100) |

Number of people that are working and not cured from treatment can also be calculated as: nn * (w / 100) * (1 - cr / 100)
Number of people that are not working and not cured from treatment can also be calculated as: nn * (nw / 100) * (1 - cr / 100)

Number of people that are working and not cured from treatment can also be calculated as: nn * (w / 100) * (1 - cr / 100)
Number of people that are not working and not cured from treatment can also be calculated as: nn * (nw / 100) * (1 - cr / 100)

Now we need to answer the important question: should a government pay for IV treatment for chronic Lyme disease? From a purely economic and financial perspective which excludes any ethical
aspect of not offering antibiotics treatment to patients that suffer from a bacterial infection that can lead to death a government should finance treatment with IV antibiotics for chronic Lyme disease if

Future government revenues [treatment] + future government cost [treatment] + treatment cost

> future government revenues [no treatment] + future government cost [no treatment]

where government revenues are positive values and government costs are negative values. Such governments’ financial chronic Lyme disease balance sheet is presented in Table 16.

| A government’s Financial Balance | Today | If the Government Finance IV Treatment | If the Government Does not Finance IV Treatment | Future | If the Government Does not Finance IV Treatment |
|----------------------------------|-------|---------------------------------------|-----------------------------------------------|--------|-----------------------------------------------|
| A government’s chronic Lyme disease revenue (+) | x1 = tax revenues from chronic Lyme patients that are cured from treatment and are working | x5 = 0 | x6 = 0 |
| A government’s chronic Lyme disease revenue (+) | x2 = saved disability benefits for chronic Lyme patients that are cured from treatment and are working | -X3 = lost tax revenues from chronic Lyme patients that are still sick after treatment and are not working | -X7 = lost tax revenues from chronic Lyme patients that are sick and have not received treatment and are not working | -x8 = disability payments to chronic Lyme patients that are still sick after treatment and are not working |
| A government’s chronic Lyme disease cost (-) | -X4 = disability payments to chronic Lyme patients that are still sick after treatment and are not working | -X5 = tax revenues from chronic Lyme patients that are cured and are working | -x22 = saved future disability benefits for chronic Lyme patients that are cured from treatment and are working | -x33 = lost future tax revenues for chronic Lyme patients that are still sick after treatment and are not working |
| A government’s chronic Lyme disease cost (-) | -X6 = saved disability benefits for chronic Lyme patients that are still sick after treatment and are not working | -x3 = tax revenues from chronic Lyme patients that are still sick after treatment and are not working | -x44 = future disability payments to chronic Lyme patients that are still sick after treatment and are not working |

Treatment cost for 0.5 year of IV antibiotics (-) - IV0.5
Treatment cost for 1 year of IV antibiotics (-) - IV1

A government’s revenues because of treatment x1 + x2 x11 + x22
A government’s costs for chronic Lyme disease with IV 0.5 year x3 + x4 + IV0.5 x33 + x44 + IV0.5
A government’s cost for chronic Lyme disease with IV 1 year x3 + x4 + IV1 x33 + x44 + IV1
A government’s cost without treatment (-) x7 + x8 x77 + x88
A government’s financial balance based on today’s revenues and costs for 0.5 year of IV treatment (GBTIV0.5) x1 + x2 + x3 + x4 + IV0.5 - x7 - x8 x11 + x22 + x33 + x44 + IV0.5 - x77 - x88
A government’s financial balance based on today’s revenues and costs for 1 year of IV treatment (GBTIV1) x1 + x2 + x3 + x4 + IV1 - x7 - x8 x11 + x22 + x33 + x44 + IV1 - x77 - x88

To justify IV treatment for 0.5 years x11 + x22 + x33 + x44 + IV0.5 > x55 + x66 + x77 + x88 x11 + x2 + x3 + x4 + IV0.5 - (x55 + x66 + x77 + x88) > 0
GBTIV0.5 = x11 + x22 + x33 + x44 + IV0.5 - x77 - x88 > 0
To justify IV treatment for 1 year x11 + x2 + x3 + x4 + IV1 - (x55 + x66 + x77 + x88) > 0
GBTIV1 = x11 + x22 + x33 + x44 + IV1 - x77 - x88 > 0

Note also that the previous equation become less reliable over time. The closer we get to our endpoint which is 2050 the less impact the future government revenues and future government...
expenditures will have on a government’s financial balance. Note that tax revenues from chronic Lyme patients that are not cured with treatment and working, saved disability benefits from chronic Lyme patients that are not cured with treatment and working, tax revenues from chronic Lyme patients that are sick and have not received treatment and are working and saved disability benefits from chronic Lyme patients that are sick and have not received treatment and are working are not included in the above government financial chronic Lyme disease balance sheet because they will be the same regardless of if the government choose to treat or not. The government’s purely financial treatment decision regarding chronic Lyme disease is further illustrated in Table 17.

Table 17. A government’s financial IV treatment decision regarding chronic Lyme disease.

| 1 | 2 | 3 | 4 | 5 | 6 |
|---|---|---|---|---|---|
| 1 = If the government finance IV treatment | 2 = If the government does not finance IV treatment | 3 = is GBFT > GBFnT ? | 4 = GBFT - GBFnT | 5 = is GFBBT - GFBnT > 0 ? | 6 = Should a government finance treatment with IV antibiotics? |
| A government’s future chronic Lyme disease revenues | GFR | GFRnT | | | |
| A government’s future chronic Lyme disease costs | - GFC | - GFCnT | | | |
| Treatment cost today for IV antibiotics | - TCT | na | | | |
| A government’s balance based on future government revenues and costs | GBFT = GFR + GFC + TCT | GBFnT = GFRnT + GFCnT | | | |
| A government’s future chronic Lyme disease revenues | 200 | 0 | | | |
| A government’s future chronic Lyme disease costs | -200 | -100 | | | |
| Treatment cost today for IV antibiotics | -500 | 100 | | | |
| A government’s balance based on future government revenues and costs | -500 | -100 | no | -400 | no | no |
| A government’s future chronic Lyme disease revenues | 200 | 0 | | | |
| A government’s future chronic Lyme disease costs | -200 | -200 | | | |
| Treatment cost today for IV antibiotics | -300 | na | | | |
| A government’s balance based on future government revenues and costs | -300 | -200 | no | -100 | no | no |
| A government’s future chronic Lyme disease revenues | 500 | 0 | | | |
| A government’s future chronic Lyme disease costs | -300 | -200 | | | |
| Treatment cost today for IV antibiotics | -200 | na | | | |
| A government’s balance based on future government revenues and costs | 0 | -200 | yes | 200 | yes | yes |
| A government’s future chronic Lyme disease revenues | 600 | 0 | | | |
| A government’s future chronic Lyme disease costs | -400 | -500 | | | |
| Treatment cost today for IV antibiotics | -100 | na | | | |
| A government’s balance based on future government revenues and costs | 100 | -500 | yes | 600 | yes | yes |

We now again assume that the annual growth rate of the infection is 2% per year, the annual population growth rate in the USA is assumed to be 1% and the annual population growth rate in Europe is assumed to be 0.2%. We assume that we treat all chronic Lyme patients with IV antibiotics for either six months or one year and we again look at the value for different variables for five different cure rates 0%, 25%, 50%, 75% and 100%. All charts in this section and more can be found in the Excel file “Lyme disease calculations.xlsx” There will be only four individual group captions in this section: today’s government costs and revenues in the USA over time, future government costs and revenues in the USA over time, today’s government costs and revenues in Europe over time and future government costs and revenues in Europe over time. There is no point in including a caption for each chart when
the same text can be found in the chart’s legend. We can see in Figure 19 that the US government’s financial balance based on today’s cost and revenues for antibiotic IV treatment for 6 months for chronic Lyme disease GBTiv0.5 is positive for an assumed cure rate of 25% and above and for 1 year of IV treatment with antibiotics GBTiv1 is positive for an assumed cure rate of 50% and above. We can see in Figure 20 that the US government’s financial balance based on future cost and revenues for antibiotic IV treatment for 6 months for chronic Lyme disease GBTiv0.5 is positive for an assumed cure rate of 25% and above and for 1 year of IV treatment GBTiv1 is also positive for an assumed cure rate of 50% and above. We can see in Figure 21 that the European governments’ financial balance based on today’s cost and revenues for antibiotic IV treatment for 6 months for chronic Lyme disease GBTiv0.5 is positive for an assumed cure rate of 25% and above and for 1 year of IV treatment with antibiotic GBTiv1 is positive for an assumed cure rate of 50% and above. We can see in Figure 22 that the European governments’ financial balance based on future cost and revenues for IV antibiotic treatment for 6 months for chronic Lyme disease GBTiv0.5 is positive for an assumed cure rate of 25% and above and for 1 year of IV antibiotic treatment GBTiv1 is also positive for an assumed cure rate of 25% and above.

Figure 19. Cont.
Figure 19. Cont.
Figure 19. Today’s government costs and revenues in the USA over time.
Figure 20. Cont.
Figure 20. Future government costs and revenues in the USA over time.
Figure 21. Cont.
Figure 21. Cont.
Figure 21. Today’s government costs and revenues in Europe over time.
Figure 22. Cont.
Figure 22. Cont.
Figure 22. Cont.
Figure 22. Future government costs and revenues in Europe over time.

4. Discussion

Independent and objective peer review is a critical component for scientific publication because of two main reasons. (1) It makes sure that only high-quality papers are published which is critical for
the credibility of a scientific journal and science in general. (2) It leads to decentralization of power which is also critical for the credibility of a scientific journal and science in general. When the CDC publishes its research on Lyme disease in a private special interest group’s “scientific” journal such as *Clinical Infectious Diseases* which is issued by the IDSA the probability that the paper will receive independent and objective peer review is close to zero. *The Journal of Infectious Diseases, Clinical Infectious Diseases* and *Open Forum Infectious Diseases* are all “scientific” journals published by the IDSA. Scientific publication is not democratic in a sense that everyone (including people without subject knowledge) can vote and where the majority decides which papers are published. Such undemocratic nature of science is a strength, but it can also sometimes be a weakness. Medicine is based on science, but medicine also has patients. Patients are the only ones that will be affected by a physician’s decision. It is therefore not unrealistic to expect that patients can influence decision making regarding their diagnosis and treatment. Unfortunately, today medical doctors, journal editors that desk reject papers, politicians, journalists and governments are experts when it comes to centralized decision-making. It also appears that medicine and medical science today has become politicized where personal and subjective opinions govern. Government health agencies have not done an objective review of the scientific literature regarding chronic Lyme disease. There exist at least five reasons for this. (1) Governments’ treatment guidelines or research on Lyme disease have not been published in high quality international scientific journals which means that such publications have not been subject to objective and independent peer review which is very unscientific. (2) Infectious disease doctors (in the USA IDSA) and government health agencies (in the USA CDC) have previously or are possibly still colluding and IDSA is colluding with insurance companies in the USA. (3) The treatment guideline panels with doctors have not been sufficiently diversified because ILADS doctors have been excluded from the decision making. (4) Chronic Lyme patients have been excluded from the treatment guideline panels for Lyme disease. (5) Governments have a fear of antibiotic resistance even though there exists no scientific evidence that the *Borrelia* bacteria have developed antibiotic resistance. Encouraging physicians to do an independent and objective evaluation of the scientific literature is more important than to force physicians to blindly follow government treatment guidelines that may or may not be ethical and up to date. Doing scientific research and reading the scientific literature are demanding work. Blindly following treatment guidelines requires little effort and does not require a degree in medicine because you don’t need to be a professional chef to follow a cooking recipe. The importance of a patient voice is also something that is missing in the debate. Doctors have a responsibility to help sick patients. To deny a patient medical treatment in a situation when there exists scientific evidence to support treatment will always be controversial. Infectious disease doctors need to be more modest here. Patients have a critical role in healthcare and without them, there will be no healthcare. Some people conclude that since oral antibiotics cannot cure chronic Lyme disease patients, oral antibiotics should not be given to chronic Lyme patients. Such reasoning is a *non-sequitur* (logical fallacy). HIV is a chronic infection that cannot be eradicated with oral medication, but HIV patients still receive oral medication for life. Does there exist a general rule in medicine that states that only viral infections can become chronic? No! Is it ethical to refuse a patient treatment that reduces a patient’s bacterial load, reduces a patient’s symptoms and sometimes even cures them? No! Is it ethical to refuse a patient treatment for an infection that can kill them without treatment? No!

There also exist at least two problems with studies that treat chronic Lyme disease patients with IV antibiotics and conclude that such treatment did not eradicate the infection is (1) These studies usually treat chronic Lyme patients with monotherapy which is not a recommended treatment. (2) They usually do not treat for a sufficient period to make inference about how effective IV treatment is in eradicating the infection. I also want to point out that a double-blind, randomized controlled trial where neither the patient nor the treating doctor knows if the patient has received placebo or medication might, in theory, sound like a good a way to determine how effective a certain medication is. Reality is however far from theory. There also exists an ethical aspect. It is not ethical to give patients with a confirmed infection placebo in the name of science when such action could result in a lifelong infection
for the patient. The estimated number of people that were infected with Lyme disease in the USA in 2008 is 829,600. If we assume that the incidence rate 2008 in the USA and Europe were the same, then approximately 2 million people were infected with Lyme disease Europe in 2008. Given an annual infection growth rate of 2% then in 2018 in the USA approximately 1 million people and in Europe 2.4 million will get infected with Lyme disease. Given an annual population growth of 1% in the USA then by 2050 in the USA 56 million people (12% of the population in the USA in 2050) will have been infected with Lyme disease. Given an annual population growth in Europe of 0.2% then by 2050 in Europe 135 million people (17% of the population Europe in 2050) will have been infected with Lyme disease. Most of these Lyme disease infections will, unfortunately, become chronic. I have also shown in this article that the financial implications of chronic Lyme disease are vast. A financial cost not only to patients but also to governments. Given that acute Lyme disease is treated with oral antibiotics for one month and chronic Lyme is treated with IV antibiotics for either 6 months or 1 year, then the estimated treatment cost for acute and chronic Lyme disease for 2018 for the USA is somewhere between 4.8 billion USD and 9.6 billion USD and for Europe somewhere between 10.1 billion EUR and 20.1 billion EUR. The estimated lost personal income for chronic Lyme patients that are not working for 2018 for the USA is 16.1 billion USD and in Europe 24.8 billion EUR. If the governments in the USA and Europe do not finance IV treatment with antibiotics for chronic Lyme disease, then the estimated government cost for chronic Lyme disease in the form of lost tax revenues because some chronic Lyme patients are not working plus disability benefits for chronic Lyme patients that not working for 2018 for the USA is 10.1 billion USD and for Europe 20.1 billion EUR. The government cost for chronic Lyme disease which does not include treatment costs for 2018 in the USA is 2.1 times larger than the cost for 6 months of IV treatment with antibiotics in the USA in 2018 and 1.1 times larger than the cost for 1 year of IV treatment with antibiotics in the USA in 2018. In Europe, the government cost for chronic Lyme disease for 2018 is two times larger than the cost for six months of IV treatment with antibiotics for Europe for 2018 and the government cost is on par with the cost, for one year of IV with antibiotics treatment for Europe for 2018. The cost for the governments of having chronic Lyme patients sick in perpetuity is very large. Future government revenues and future government costs for chronic Lyme disease are however much more important to look at than historical annual cost. Governments aggressive way to try to reduce antibiotic use unfortunately only creates more chronic infections where people can be forced to take oral antibiotics for the rest of their lives which means that the initial reduction in antibiotics use that can be observed due to the government’s aggressive antibiotics reduction policy increases the use of antibiotics in the long run. No one thinks that diseases that are caused by a virus should be treated with antibiotics but when people who need antibiotics for their survival do not get access to them, then there is something wrong.

5. Conclusions

If someone acknowledges that acute Lyme disease is caused by bacteria, then they are also forced by logical reasoning to acknowledge that chronic Lyme disease is caused by bacteria because the immune system by itself can never eradicate a Borrelia infection and a bacterial infection does not magically disappear without antibiotic treatment once a person has been infected. We have known for a long time that antibiotics kill bacteria. However, sometimes oral and even IV antibiotics are unable to eradicate a Borrelia infection, but those observations are irrelevant because HIV patients receive treatment for life even though antiviral medications cannot eradicate an HIV infection and people with chronic Lyme disease die without antibiotic treatment. One single death caused by chronic Lyme disease is also enough to justify antibiotic treatment for chronic Lyme disease because a killer must “only” kill one person to become a killer. When a private special interest group such as the IDSA makes a political claim that three weeks of oral antibiotics is always enough to eradicate an acute or chronic Lyme disease infection, then the burden of proof lies with the IDSA. The IDSA must show the scientific evidence that supports their claim. I can claim that a red car is circulating the moon but unless I can show the scientific evidence that supports my claim, my claim has little meaning.
Lyme disease and chronic Lyme disease are well-hidden pandemics because: (1) the tests for Lyme disease are insensitive. Even if patients are “lucky” enough to test positive, patients will sometimes not receive antibiotic treatment because then the infectious disease physicians (in the USA IDSA) claim without scientific evidence that the antibodies are old and not a sign of an active infection. It becomes impossible for a patient to win. (2) the IDSA and CDC have previously or are possibly still colluding. (3) Because the cost of IV antibiotic treatment for chronic Lyme disease is so large insurance companies in the USA collude with IDSA to make sure IDSA deny that chronic Lyme disease is real disease that is caused by bacteria and require antibiotic treatment. This leads to that many Lyme disease patients are denied treatment and left to suffer until the infection eventually kills them. For many chronic Lyme disease patients, ILADS doctors, therefore, become heroes because these medical doctors are treating extremely sick and abandoned patients with long-term antibiotics. The CDC reported that in 2008 in the USA 444,000 people (high scenario) were infected with Lyme disease. The CDC assumption, without motivation or empirical evidence, that the number of healthy people in the control group is 4.4 times the number of infected people in the disease group for the high scenario is not realistic. If the CDC had assumed that the size of the control group with healthy people was the same size as the disease group with infected people ($p = 1$), then the number of Lyme disease infections in the USA for 2008 for the CDC high scenario would have been 527,288 instead of 444,000 given a Lyme disease test sensitivity of 0.669 and specificity of 0.961. If we assume a 1 to 1 relationship between the two groups and a Lyme disease test sensitivity of 0.44 and a Lyme disease test specificity of 0.99, then we get 829,600 infections in the USA in 2008. Because there exists uncertainty if the CDC is independent and objective, there also exists uncertainty about the CDC ability to protect public health. This research paper is one of the first studies that have estimated the financial cost of chronic Lyme disease both for individuals and governments. This research paper is also the first paper that can show that if the USA and European governments want to minimize future cost and maximize future revenues, then they should pay for IV treatment with antibiotics for chronic Lyme disease up to a year even if the estimated cure rate is as low as 25%. I am convinced that the history books in the future will describe controversy that exists today regarding chronic Lyme disease as one of the most shameful affairs in medicine.

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**VBA Code**

```
Public Sub zzz()
  se = 0.44
  sp = 0.99
  np = 26
  nx = 10

  ThisWorkbook.Worksheets("zzz sub procedure").Range("B2:np" & np + 1).NumberFormat = "0.00"
  ThisWorkbook.Worksheets("zzz sub procedure").Range("A1:np" & np + 1).HorizontalAlignment = xlCenter

  Dim x As Long
  Dim y As Long
  Dim i As Integer
  Dim j As Integer
  Dim pp()
  Dim xx()
  Dim m()
```
For i = 1 To np
    For j = 1 To nx
        ReDim Preserve pp(i)
        ReDim Preserve xx(j)
        ReDim m(1 To np, 1 To nx)
        pp(i) = (i - 1) / 5
        xx(j) = j * 10
        m(i, j) = ((xx(j) * se + pp(i) * xx(j) * (1 - sp)) / (xx(j) + pp(i) * xx(j))) * 100
    Next j
Next i

End Sub

Public Function Lyme1(se, sp, p, z, Output As String)
    Dim A(1 To 2, 1 To 2)
    Dim C(1 To 2, 1 To 1) As Long
    Dim x As Long
    Dim y As Long
    A(1, 1) = se
    A(1, 2) = 1 - sp
    A(2, 1) = -p
    A(2, 2) = 1
    C(1, 1) = z
    C(2, 1) = 0
    r = WorksheetFunction.MMult(WorksheetFunction.MInverse(A), C)
    x = r(1, 1)
    y = r(2, 1)
    zz = (z / (x + y)) * 100
    If Output = “x” Then Lyme1 = x
    If Output = “y” Then Lyme1 = y
    If Output = “zz” Then Lyme1 = zz
End Function

Public Function Lyme2(se, sp, p, z, Output As String)
    Dim x As Long
Dim y As Long

x = z / (-sp * p + se + p)
y = p * z / (-sp * p + se + p)
zz = (z / (x + y)) * 100

If Output = “x” Then Lyme2 = x
If Output = “y” Then Lyme2 = y
If Output = “zz” Then Lyme2 = zz

End Function

Public Function Lyme3(se, sp, p, z, zz, Output As String)

Dim x As Long
Dim y As Long

x = 100 * z / (p * zz + zz)
y = 100 * p * z / (p * zz + zz)

If Output = “x” Then Lyme3 = x
If Output = “y” Then Lyme3 = y

End Function

References

1. Harvey, W.T.; Salvato, P. “Lyme disease”: Ancient engine of an unrecognized borreliosis pandemic? *Med. Hypotheses* **2003**, *60*, 742–759. [CrossRef]
2. Cameron, D.J. Proof That Chronic Lyme Disease Exists. *Interdiscip. Perspect. Infect. Dis.* **2010**. Available online: [http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2876246/](http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2876246/) (accessed on 21 March 2017).
3. ECDC. Borreliosis. *Borre.* 2017. Available online: [http://ecdc.europa.eu/en/healthtopics/emerging_and_vector-borne_diseases/tick_borne_diseases/lyme_disease/pages/index.aspx](http://ecdc.europa.eu/en/healthtopics/emerging_and_vector-borne_diseases/tick_borne_diseases/lyme_disease/pages/index.aspx) (accessed on 21 March 2017).
4. Tonks, A. Lyme wars. *BMJ* **2007**, *335*, 910–912. [CrossRef] [PubMed]
5. Stricker, R.B.; Johnson, L. Lyme Disease: Call for a “Manhattan Project” to Combat the Epidemic. *PLoS Pathog.* **2014**, *10*, e1003796. [CrossRef] [PubMed]
6. IDSA. IDSA: Infectious Diseases Society of America. Available online: [http://www.idsociety.org/Index.aspx](http://www.idsociety.org/Index.aspx) (accessed on 8 December 2017).
7. ILADS. ILADS—End Lyme disease epidemic through education, awareness & physician training. Available online: [http://www.ilads.org/](http://www.ilads.org/) (accessed on 8 December 2017).
8. Wormser, G.P.; Dattwyler, R.J.; Shapiro, E.D.; Halperin, J.J.; Steere, A.C.; Klempner, M.S.; Krause, P.J.; Bakken, J.S.; Strie, F.; Stanek, G.; et al. The Clinical Assessment, Treatment, and Prevention of Lyme Disease, Human Granulocytic Anaplasmosis, and Babesiosis: Clinical Practice Guidelines by the Infectious Diseases Society of America. *Clin. Infect. Dis.* **2006**, *43*, 1089–1134. [CrossRef] [PubMed]
9. Donta, S.T. Issues in the diagnosis and treatment of lyme disease. *Open Neurol. J.* **2012**, *6*, 140–145. [CrossRef] [PubMed]
10. Sharma, B.; Brown, A.V.; Matluck, N.E.; Hu, L.T.; Lewis, K. Borrelia burgdorferi, the Causative Agent of Lyme Disease, Forms Drug-Tolerant Persister Cells. *Antimicrob. Agents Chemother.* **2015**, *59*, 4616–4624. [CrossRef] [PubMed]
11. Caskey, J.R.; Embers, M.E. Persister Development by Borrelia burgdorferi Populations In Vitro. *Antimicrob. Agents Chemother.* **2015**, *59*, 6288–6295. [CrossRef] [PubMed]
12. Hodzic, E.; Imai, D.; Feng, S.; Barthold, S.W. Resurgence of Persisting Non-Cultivable Borrelia burgdorferi following Antibiotic Treatment in Mice. *PLoS ONE* **2014**, *9*, e86907. [CrossRef] [PubMed]
13. Hodzic, E.; Feng, S.; Holden, K.; Freet, K.J.; Barthold, S.W. Persistence of Borrelia burgdorferi following antibiotic treatment in mice. *Antimicrob. Agents Chemother.* 2008, 52, 1728–1736. [CrossRef] [PubMed]

14. Barthold, S.W.; Hodzic, E.; Imai, D.M.; Feng, S.; Yang, X.; Luft, B.J. Ineffectiveness of Tigecycline against Persistent Borrelia burgdorferi. *Antimicrob. Agents Chemother.* 2010, 54, 643–651. [CrossRef] [PubMed]

15. Yrjänäinen, H.; Hytönen, J.; Song, X.R.; Oksi, J.; Hartiala, K.; Viljanen, M.K. Anti-tumor necrosis factor-alpha treatment activates Borrelia burgdorferi spirochetes 4 weeks after ceftriaxone treatment in C3H/He mice. *J. Infect. Dis.* 2007, 195, 1489–1496. [CrossRef] [PubMed]

16. Straubinger, R.K.; Summers, B.A.; Chang, Y.F.; Appel, M.J. Persistence of Borrelia burgdorferi in experimentally infected dogs after antibiotic treatment. *J. Clin. Microbiol.* 1997, 35, 111–116. [PubMed]

17. Chang, Y.-F.; Ku, Y.-W.; Chang, C.-F.; Chang, C.-D.; McDonough, S.P.; Divers, T.; Pough, M. Antibiotic treatment of experimentally Borrelia burgdorferi-infected ponies. *Vet. Microbiol.* 2005, 107, 285–294. [CrossRef] [PubMed]

18. Embers, M.E.; Barthold, S.W.; Borda, J.T.; Bowers, L.; Doyle, L.; Hodzic, E.; Jacobs, M.B.; Hasenkampf, N.R.; Martin, D.S.; Narasimhan, S.; et al. Persistence of Borrelia burgdorferi in Rhesus Macaques following Antibiotic Treatment of Disseminated Infection. *PLoS ONE* 2012, 7, e29914. [CrossRef]

19. Rudenko, N.; Golovchenko, M.; Vancova, M.; Clark, K.; Grubhoffer, L.; Oliver, J.H. Isolation of live Borrelia burgdorferi from patients with chronic Lyme disease symptoms—A PCR study of 97 cases. *Neurology* 1998, 51, 1489–1491. [CrossRef] [PubMed]

20. Phillips, S.E.; Mattman, L.H.; Hulinská, D.; Moayad, H. A proposal for the reliable culture of Borrelia burgdorferi from patients with chronic Lyme disease, even from those previously aggressively treated. *Infection* 1999, 26, 364–367. [CrossRef] [PubMed]

21. Schmidli, J.; Hunziker, T.; Moesi, P.; Schaad, U.B. Cultivation of Borrelia burgdorferi from Joint Fluid Three Months after Treatment of Facial Palsy Due to Lyme Borreliosis. *J. Infect. Dis.* 1988, 158, 905–906. [CrossRef] [PubMed]

22. Cimmino, M.A.; Azzolini, A.; Tobia, F.; Pesce, C.M. Spirochetes in the spleen of a patient with chronic Lyme disease. *Am. J. Clin. Pathol.* 1989, 91, 95–97. [CrossRef] [PubMed]

23. Treib, J.; Fernandez, A.; Haass, A.; Grauer, M.; Holzer, G.; Woessner, R. Clinical and serologic follow-up in patients with neuroborreliosis. *Neurology* 1998, 51, 1489–1491. [CrossRef] [PubMed]

24. Preac-Mursic, V.; Weber, K.; Pfister, H.W.; Wilske, B.; Gross, B.; Baumann, A.; Prokop, J. Survival of Borrelia burgdorferi in antibiotically treated patients with Lyme borreliosis. *Infection* 1999, 27, 355–359. [CrossRef] [PubMed]

25. Hudson, B.J.; Stewart, M.; Lennox, V.A.; Fukunaga, M.; Yabuki, M.; Macorison, H.; Kitchener-Smith, J. Culture-positive Lyme borreliosis. *Med. J. Aust.* 1998, 168, 500–502. [PubMed]

26. Kirsch, M.; Ruben, F.L.; Steere, A.C.; Duray, P.H.; Norden, C.W.; Winkelstein, A. Fatal Adult Respiratory Distress Syndrome in a Patient With Lyme Disease. *JAMA* 1988, 259, 2737–2739. [CrossRef] [PubMed]

27. Valesová, H.; Maier, J.; Havlík, J.; Hulinská, D.; Hercogová, J. Long-term results in patients with Lyme arthritis following treatment with ceftriaxone. *Infection* 1996, 24, 98–102. [PubMed]

28. Asch, E.S.; Bujak, D.I.; Weiss, M.; Peterson, M.G.; Weinstein, A. Lyme disease: an infectious and postinfectious Distress Syndrome in a Patient With Lyme Borreliosis. *J. Rheumatol.* 1994, 21, 454–461. [PubMed]

29. Pfister, H.W.; Preac-Mursic, V.; Wilske, B.; Schielke, E.; Sörgel, F.; Einhäupl, K.M. Randomized comparison of ceftriaxone and cefotaxime in Lyme neuroborreliosis. *J. Infect. Dis.* 1991, 163, 311–318. [CrossRef] [PubMed]

30. Breier, F.; Khanakah, G.; Stanek, G.; Kunz, G.; Aberer, E.; Schmidt, B.; Tappeiner, G. Isolation and polymerase chain reaction typing of Borrelia afzelii from a skin lesion in a seronegative patient with generalized ulcerating bullous lichen sclerosus et atrophicus. *Br. J. Dermatol.* 2001, 144, 387–392. [CrossRef] [PubMed]

31. Oksi, J.; Marjamäki, M.; Nikoskelainen, J.; Viljanen, M.K. Borrelia burgdorferi detected by culture and PCR in clinical relapse of disseminated Lyme borreliosis. *Ann. Med.* 1999, 31, 225–232. [CrossRef] [PubMed]

32. Bayer, M.E.; Zhang, L.; Mayer, M.H. Borrelia burgdorferi DNA in the urine of treated patients with chronic Lyme disease symptoms—A PCR study of 97 cases. *Infection* 1996, 24, 347–353. [CrossRef] [PubMed]

33. Battafarano, D.F.; Combs, J.A.; Enzenauer, R.J.; Fitzpatrick, J.E. Chronic septic arthritis caused by Borrelia burgdorferi. *Clin. Orthop. Relat. Res.* 1993, 297, 238–241.
34. Häupl, T.; Hahn, G.; Rittig, M.; Krause, A.; Schoerner, C.; Schönherr, U.; Kalden, J.R.; Burmester, G.R. Persistence of Borrelia burgdorferi in ligamentous tissue from a patient with chronic Lyme borreliosis. *Arthritis Rheumatol.* 1993, 36, 1621–1626. [CrossRef]

35. Preac-Mursic, V.; Pfister, H.W.; Spiegel, H.; Burk, R.; Wilske, B.; Reinhardt, S.; Böhmer, R. First isolation of Borrelia burgdorferi from an iris biopsy. *J. Clin. Neuroophthalmol.* 1993, 13, 155–161. [PubMed]

36. Lawrence, C.; Lipton, R.B.; Lowy, F.D.; Coyle, P.K. Seronegative chronic relapsing neuroborreliosis. *Eur. Neurol.* 1995, 35, 113–117. [CrossRef] [PubMed]

37. Johnson, L.; Stricker, R.B. Treatment of Lyme disease: A medicolegal assessment. *Expert Rev. Anti-Infect. Ther.* 2004, 2, 533–557. [CrossRef] [PubMed]

38. Stricker, R.B.; Johnson, L. Persistent infection in chronic lyme disease: Does form matter? *Res. J. Infect. Dis.* 2013, 1, 2.

39. Stricker, R.B.; Johnson, L. Lyme disease: The next decade. *Infect. Drug Resist.* 2011, 4, 1–9. [CrossRef] [PubMed]

40. Melchers, W.; Meis, J.; Rosa, P.; Claas, E.; Nohlmans, L.; Koopman, R.; Horrevorts, A.; Galama, J. Amplification of Borrelia burgdorferi DNA in skin biopsies from patients with Lyme disease. *J. Clin. Microbiol.* 1991, 29, 2401–2406. [PubMed]

41. Pachner, A.R.; Cadavid, D.; Shu, G.; Dail, D.; Pachner, S.; Hodzic, E.; Barthold, S.W. Central and peripheral nervous system infection, immunity, and inflammation in the NHP model of Lyme borreliosis. *Ann. Neurol.* 2001, 50, 330–338. [CrossRef] [PubMed]

42. Goodman, J.L.; Jurkovich, P.; Kodner, C.; Johnson, R.C. Persistent cardiac and urinary tract infections with Borrelia burgdorferi in experimentally infected Syrian hamsters. *J. Clin. Microbiol.* 1991, 29, 894–896. [PubMed]

43. Steere, A.C.; Coburn, J.; Glickstein, L. The emergence of Lyme disease. *J. Clin. Invest.* 2004, 113, 1093–1101. [PubMed]

44. Berndtson, K. Review of evidence for immune evasion and persistent infection in Lyme disease. *Int. J. Gen. Med.* 2013, 6, 291–306. [CrossRef] [PubMed]

45. Adrion, E.R.; Aucott, J.; Lemke, K.W.; Weiner, J.P. Health Care Costs, Utilization and Patterns of Care following Lyme Disease. *PLoS ONE* 2015, 10, e0116767. [CrossRef] [PubMed]

46. Stricker, R.B.; Johnson, L. Lyme disease: The promise of Big Data, companion diagnostics and precision medicine. *Infect. Drug Resist.* 2016, 9, 215–219. [CrossRef] [PubMed]

47. IDSA. IDSA: Updated Guidelines on Diagnosis, Treatment of Lyme Disease. Available online: http://www.idsociety.org/Updated_Guidelines_on_Diagnosis_Treatment_of_Lyme_Disease/ (accessed on 8 December 2017).

48. Maloney, E.L. Controversies in Persistent (Chronic) Lyme Disease. *J. Infus. Nurs.* 2016, 39, 369–375. [CrossRef] [PubMed]

49. Johnson, L.; Stricker, R.B. Attorney General forces Infectious Diseases Society of America to redo Lyme guidelines due to flawed development process. *J. Med. Ethics.* 2009, 35, 283–288. [CrossRef] [PubMed]

50. Johnson, L.; Stricker, R.B. The Infectious Diseases Society of America Lyme guidelines: A cautionary tale about the development of clinical practice guidelines. *Philos. Ethics Hum. Med.* 2010, 5, 9. [CrossRef] [PubMed]

51. CourtDocumentLymeDisease—11 October. 2017. Available online: https://www.courthousenews.com/wp-content/uploads/2017/11/LymeDisease.pdf (accessed on 8 December 2017).

52. Langford, C. Insurers Accused of Conspiring to Deny Lyme Disease Coverage. 2017. Available online: https://www.courthousenews.com/insurers-accused-conspiring-deny-lyme-disease-coverage/ (accessed on 8 December 2017).

53. Multiple Chronic Conditions Chartbook. 2010. Available online: https://www.ahrq.gov/sites/default/files/wysiwyg/professionals/prevention-chronic-care/decision/mcc/mccchartbook.pdf (accessed on 9 December 2017).

54. Johnson, L.; Stricker, R.B. Final Report of the Lyme Disease Review Panel of the Infectious Diseases Society of America: A Pyrrhic Victory? *Clin. Infect. Dis.* 2010, 51, 1108–1109. [CrossRef] [PubMed]
55. The National Academies of Sciences, Engineering, and Medicine. Standards for Developing Trustworthy Clinical Practice Guidelines: Health and Medicine Division. 2011. Available online: http://www.nationalacademies.org/hmd/Reports/2011/Clinical-Practice-Guidelines-We-Can-Trust/Standards.aspx (accessed on 17 March 2017).

56. Lee, D.H.; Vielemeyer, O. Analysis of Overall Level of Evidence behind Infectious Diseases Society of America Practice Guidelines. *Arch. Intern. Med.* 2011, 171, 18–22. [CrossRef] [PubMed]

57. CDC: Lyme carditis. What you need to know about Lyme carditis. 2016. Available online: https://www.cdc.gov/lyme/signs_symptoms/lymecarditis.html (accessed on 17 March 2017).

58. Stricker, R.B.; Delong, A.K.; Green, C.L.; Savely, V.R.; Chamallas, S.N.; Johnson, L. Benefit of intravenous antibiotic therapy in patients referred for treatment of neurologic Lyme disease. *Int. J. Gen. Med.* 2011, 4, 639–646. [CrossRef] [PubMed]

59. Thyresson, N. The penicillin treatment of acrodermatitis chronica (Herxheimer). *Acta Derm. Venereol.* 1949, 29, 572–621. [PubMed]
78. Keller, A.; Graefen, A.; Ball, M.; Matzas, M.; Boisguerin, V.; Maixner, F.; Leidinger, P.; Backes, C.; Khairat, R.; et al. New insights into the Tyrolean Iceman’s origin and phenotype as inferred by whole-genome sequencing. *Nat. Commun.* 2012, 3, 698. [CrossRef] [PubMed]

79. Poinar, G. Spirochete-like cells in a Dominican amber Amblyomma tick (Arachnida: Ixodidae). *Hist. Biol.* 2015, 27, 565–570. [CrossRef]

80. Fraser, C.M.; Casjens, S.; Huang, W.M.; Sutton, G.G.; Clayton, R.; Lathigra, R.; White, O.; Ketchum, K.A.; Dodson, R.; Hickey, E.K.; et al. Genomic sequence of a Lyme disease spirochaete, *Borrelia burgdorferi*. *Nature* 1997, 390, 580–586. [CrossRef]

81. Di, L.; Pagan, P.E.; Packer, D.; Martin, C.L.; Akther, S.; Ramrattan, G.; Mongodin, E.F.; Fraser, C.M.; Schutzer, S.E.; Luft, B.J.; et al. BorreliaBase: A phylogeny-centered browser of Borrelia genomes. *BMC Bioinform.* 2014, 15, 233. [CrossRef] [PubMed]

82. Bunikis, J.; Garpmo, U.; Tsao, J.; Berglund, J.; Fish, D.; Barbour, A.G. Sequence typing reveals extensive strain diversity of the Lyme borreliosis agents *Borrelia burgdorferi* in North America and *Borrelia afzelii* in Europe. *Microbiology* 2004, 150, 1741–1755. [CrossRef] [PubMed]

83. Rudenko, N.; Golovchenko, M.; Grubhoffer, L.; Oliver, J.H. *Borrelia carolinensis* sp. nov., a New (14th) Member of the Borrelia burgdorferi Sensu Lato Complex from the Southeastern Region of the United States. *J. Clin. Microbiol.* 2009, 47, 134–141. [CrossRef] [PubMed]

84. Wilske, B.; Barbour, A.G.; Bergström, S.; Burman, N.; Restrepo, B.I.; Rosa, P.A.; Schwann, T.; Soutschek, E.; Wallich, R. Antigenic variation and strain heterogeneity in *Borrelia* spp. *Res. Microbiol.* 1992, 143, 583–596. [CrossRef]

85. Baranton, G.; Seinost, G.; Theodore, G.; Postic, D.; Dykhuizen, D. Distinct levels of genetic diversity of *Borrelia burgdorferi* are associated with different aspects of pathogenicity. *Res. Microbiol.* 2001, 152, 149–156. [CrossRef]

86. Nardelli, D.T.; Callister, S.M.; Schell, R.F. Lyme Arthritis: Current Concepts and a Change in Paradigm. *Clin. Vaccine Immunol.* 2008, 15, 21–34. [CrossRef] [PubMed]

87. Sperling, J.; Middelveen, M.; Klein, D.; Sperling, F. Evolving Perspectives on Lyme Borreliosis in Canada. *Open Neurol. J.* 2012, 6, 94–103. [CrossRef] [PubMed]

88. Zhang, X.-C.; Yang, Z.-N.; Lu, B.; Ma, X.-F.; Zhang, C.-X.; Xu, H.-J. The composition and transmission of microbiome in hard tick, *Ixodes persulcatus*, during blood meal. *Ticks Tick Borne Dis.* 2014, 5, 864–870. [CrossRef] [PubMed]

91. Brorson, O.; Brorson, S.H. Transformation of cystic forms of *Borrelia burgdorferi* to normal, mobile spirochetes. *Infection* 1997, 25, 240–246. [CrossRef] [PubMed]

92. Meriläinen, L.; Herranen, A.; Schwarzbach, A.; Gilbert, L. Morphological and biochemical features of *Borrelia burgdorferi* pleomorphic forms. *Microbiology* 2015, 161, 516–527. [CrossRef] [PubMed]

93. Sapi, E.; Bastian, S.L.; Mpoy, C.M.; Scott, S.; Rattelle, A.; Pabbiati, N.; Poruri, A.; Burugu, D.; Theophilus, P.A.S.; Pham, T.V.; et al. Characterization of Biofilm Formation by *Borrelia burgdorferi* In Vitro. *PLoS ONE* 2012, 7, e48277. [CrossRef] [PubMed]

94. Smith, A.J.; Oertle, J.; Prato, D. Chronic Lyme Disease: Persistent Clinical Symptoms Related to Immune Evasion, Antibiotic Resistance and Various Defense Mechanisms of *Borrelia burgdorferi*. *Open J. Med. Microbiol.* 2014, 4, 252. [CrossRef]

95. Zhang, J.-R.; Hardham, J.M.; Barbour, A.G.; Norris, S.J. Antigenic Variation in Lyme Disease *Borrelia*ae by Promiscuous Recombination of VMP-like Sequence Cassettes. *Cell* 1997, 89, 275–285. [CrossRef]

96. Liang, F.T.; Yan, J.; Mbow, M.L.; Sviat, S.L.; Gilmore, R.D.; Mamula, M.; Fikrig, E. *Borrelia burgdorferi* Changes Its Surface Antigenic Expression in Response to Host Immune Responses. *Infect. Immun.* 2004, 72, 5759–5767. [CrossRef] [PubMed]

97. Coutte, L.; Botkin, D.J.; Gao, L.; Norris, S.J. Detailed Analysis of Sequence Changes Occurring during vlsE Antigenic Variation in the Mouse Model of *Borrelia burgdorferi* Infection. *PLoS Pathog.* 2009, 5, e1000293. [CrossRef] [PubMed]
123. Brorson, O.; Brorson, S.H. In vitro conversion of Borrelia burgdorferi to cystic forms in spinal fluid, and transformation to mobile spirochetes by incubation in BSK-H medium. *Infection* 1998, 26, 144–150. [CrossRef] [PubMed]

124. Schoonjans, F. Diagnostic test evaluation calculator. *MedCalc*. 2017. Available online: https://www.medcalc.org/calc/diagnostic_test.php (accessed on 18 March 2017).

125. Middelveen, M.J.; Burke, J.; Sapi, E.; Bandoski, C.; Filush, K.R.; Wang, Y.; Franco, A.; Timmaraju, A.; Schlinger, H.A.; Mayne, P.; et al. 2015. Available online: http://f1000research.com/articles/3-309/v3 (accessed on 18 March 2017).

126. Estrada-Peña, A.; de la Fuente, J. The ecology of ticks and epidemiology of tick-borne viral diseases. *Antivir. Res.* 2014, 108, 104–128. [CrossRef] [PubMed]

127. CDC HIV. *HIV Transmission Rates in the United States, 1977–2006. Dramatic Declines Indicate Success in HIV Prevention Nationwide—CDC_TransmissionRates_FactSheet.pdf*. 2006. Available online: https://www.cdc.gov/nchhstp/newsroom/docs/CDC_TransmissionRates_FactSheet.pdf (accessed on 19 March 2017).

128. Hinckley, A.F.; Connally, N.P.; Meek, J.I.; Johnson, B.J.; Kemperman, M.M.; Feldman, K.A.; White, J.L.; Mead, P. Lyme Disease Testing by Large Commercial Laboratories in the United States. *Clin. Infect. Dis.* 2014, 59, 676–681. [CrossRef] [PubMed]

129. Hinckley, A.F. Supplementary data to Lyme disease testing by large commercial laboratories in the United States. 2014. Available online: https://oup.silverchair-cdn.com/oup/backfile/Content_public/Journal/id/39/5/10.1093_cid_ciu397/1/ciu397_Supplementary_Data.zip?Expires=1512837512&Signature=RsW8VKdIBX6Gb5tq6E3G-VQQomLIJ130MhRvsvWxGLvavuly1rzq9gAA3EYqBqipqeBrvexpqJxJwLk57DNd0IQFZ-3pnm9t8W7f1d21NQq5wv-7fb0lpoOMfITE9GFgjHFP-XxbWJyb4VORXb-xy1yvajlg1ctn3OViArLAIeAg5yCnUllZkWhbmtSmljRnlEl-CZ7lXeVjFcmPi1w4IcZ57Dx791E13wH-1KDx06nC8t7LPcphsbZkcHEVLmWw6ad1Rf1T7DswfLrlmdqPjWuj1CiZ7lhxvJfpyEO4VkdvGLysbo1dfjS5Gmo0qjkjv_&Key-Pair-Id=AIFKAliUZBIA4LVPAW3Q (accessed on 10 September 2017).

130. Seemann, E. Are Social Security Disability Benefits Taxable? 2015. Available online: https://www.labovick.com/2015/02/13/social-security-disability-benefits-taxable/ (accessed on 17 October 2017).

131. Sweden’s government tax agency. Skatteregler för privatpersoner SKV 330 utgåva 38. 2017. Available online: https://www.skatteverket.se/download/18.57cadbbd15a3688f448d20/1488271841075/skatteregler-for-privatpersoner-skv330-utgava38.pdf (accessed on 10 September 2017).

132. Johnson, L.; Wilcox, S.; Mankoff, J.; Stricker, R.B. Severity of chronic Lyme disease compared to other chronic conditions: A quality of life survey. *PeerJ* 2014, 2, e322. [CrossRef] [PubMed]

133. OECD Statistics. OECD. Available online: http://stats.oecd.org/index.aspx?r=659459&errocode=403&lastaction=login_submit (accessed on 17 October 2017).

134. TKPMG. Individual income tax rates table. 2016. Available online: https://home.kpmg.com/xx/en/home/services/tax/tax-tools-and-resources/tax-rates-online/individual-income-tax-rates-table.html (accessed on 17 October 2017).

135. Disability Secrets, B. How Much in Social Security Disability Benefits Can You Get? Disability Secrets. 2016. Available online: http://www.disabilitysecrets.com/how-much-in-ssd.html (accessed on 18 March 2017).

136. Swedish government. Sjukersättning vid Sjukdom. Available online: https://www.forsakringskassan.se/privatpers/sjuk/sjuk_minst_1_ar/sjukersattning (accessed on 17 October 2017).