Prospective comparison of two enzyme-linked immunosorbent spot assays for the diagnosis of Lyme neuroborreliosis

T. van Gorkom,*‡ W. Voet,‡ S. U. C. Sankatsing,§ C. D. M. Nijhuis,‡ E. ter Haak,† K. Kremer† and S. F. T. Thijsen*
*Department of Medical Microbiology and Immunology, Diakonessenhuis Hospital, Utrecht, the Netherlands, ‡Centre for Infectious Diseases Research, Diagnostics and Laboratory Surveillance, Centre for Infectious Disease Control, National Institute for Public Health and the Environment (RIVM), Bilthoven, §Department of Neurology, Diakonessenhuis Hospital, and †Department of Internal Medicine, Diakonessenhuis Hospital, Utrecht.

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Correspondence: S. F. T. Thijsen, Department of Medical Microbiology and Immunology, Diakonessenhuis Hospital, Bosboomstraat 1, 3508 TG Utrecht, the Netherlands.
E-mail: sthijsen@diakhuis.nl

Summary
Commercial cellular tests are used to diagnose Lyme borreliosis (LB), but studies on their clinical validation are lacking. This study evaluated the utility of an in-house and a commercial enzyme-linked immunosorbent spot (ELISpot) assay for the diagnosis of Lyme neuroborreliosis (LNB). Prospectively, peripheral blood mononuclear cells (PBMCs) were isolated from patients and controls and analysed using an in-house Borrelia ELISpot assay and the commercial LymeSpot assay. B. burgdorferi B31 whole cell lysate and a mixture of outer surface proteins were used to stimulate the PBMCs and the numbers of interferon-gamma-secreting T cells were measured. Results were evaluated using receiver operating characteristic (ROC) curve analysis. Eighteen active and 12 treated LNB patients, 10 healthy individuals treated for an early (mostly cutaneous) manifestation of LB in the past and 47 untreated healthy individuals were included. Both assays showed a poor diagnostic performance with sensitivities, specificities, positive and negative predictive values ranging from 44.4–66.7%, 42.0–72.5%, 21.8–33.3% and 80.5–87.0%, respectively. The LymeSpot assay performed equally poorly when the calculation method of the manufacturer was used. Both the in-house and the LymeSpot assay are unable to diagnose active LNB or to monitor antibiotic treatment success.

Keywords: Borrelia, Lyme neuroborreliosis, ELISpot, T cells, interferon-gamma

Introduction
Lyme borreliosis (LB) is a tick-borne disease caused by bacteria of the Borrelia burgdorferi sensu latu group. In Europe, the most prevalent species that cause LB are B. afzelii, B. garinii and B. burgdorferi sensu stricto. The most common manifestation of LB is erythema migrans (EM); other manifestations include Lyme neuroborreliosis (LNB), Lyme arthritis (LA) and acrodermatitis chronica atrophicans (ACA). Most people, however, do not notice any symptoms and clear the infection unknowingly. In the Netherlands, surveys among general practitioners conducted in 1994 and 2017 showed a fourfold increase from an estimated 6500 to an estimated 25 500 patients with EM [1,2]. In addition, 1500 cases of a disseminated manifestation of LB were reported in 2017 [1]. Thus, LB has an increasing impact on public health in the Netherlands [3].

The diagnosis of LB depends on clinical symptoms and can sometimes be difficult due to the lack of a ‘gold standard’ test, such as culture or polymerase chain reaction (PCR). Culture is only useful for skin manifestations such as EM or ACA, but is not recommended because of the varying sensitivity and long duration, and EM is mainly a clinical diagnosis [4–6]. PCR is particularly useful in skin manifestations and LA [5,7]. For LNB, both culture and PCR show varying sensitivity and are mostly useful in the early phase of the disease [5,7–10].

The most frequently used laboratory test for LB is based on the detection of Borrelia-specific antibodies. Unfortunately, the interpretation of serological tests can be difficult, as Borrelia-specific antibodies can persist lifelong and, hence, do not discriminate between an active LB and a cleared infection. Furthermore, the absence of Borrelia-specific antibodies in the early phase of the infection does not exclude LB [11]. Therefore, better diagnostic tools are needed that can establish an active LB, especially because early antibiotic therapy has proved to be effective [12].
In recent years, various cellular assays for the diagnosis of LB have been described. Some of these assays are based on the proliferation of T cells, such as the lymphocyte transformation test (LTT) described by von Baehr et al. [13] or the LTT-memory lymphocyte immunostimulation assay (MELISA) described by Valentine-Thon et al. [14]. Other assays detect cytokines which are secreted by T cells upon stimulation with *Borrelia* antigens, such as the Quantiferon test described by Callister et al. [15] or the enzym-linked immunosorbent spot (ELISpot) assay (iSpot Lyme) described by Jin et al. [16]. Most studies on cellular assays have used poorly described study populations and lack clinical validations. Despite the lack of such validations, these assays are used in some laboratories for the diagnosis of LB [17–19], and when the test result is positive – thus when *Borrelia*-specific T cells are detected – (long-term) antibiotic treatment regimes are started for treatment of active LB [17], which is of major concern. Therefore, we recently validated an in-house *Borrelia* ELISpot assay for the detection of active LNB on a well-established study population of active LNB patients and various control groups [20]. We concluded that the T-cell activity measured in our in-house *Borrelia* ELISpot assay could not be used as a marker for active LNB. In the current study, we evaluated the diagnostic performance of a commercial LymeSpot assay that has not been validated previously, and compared this to the diagnostic performance of our in-house *Borrelia* ELISpot assay in patients suspected of LNB.

**Materials and methods**

**Study population**

Inclusion for this study started in March 2014 and ended in November 2017, and for a large part ran in parallel with two previously published studies [20,21]. Therefore, most of the study participants in the current study also participated in the previous studies and, hence, the study groups of this study consisted of subgroups of the study groups of these previous studies.

All patients diagnosed with LNB in the Diakonessenhuis Hospital, Utrecht, the St Antonius Hospital, Nieuwegein and the National Institute for Public Health and the Environment (RIVM), Bilthoven, the Netherlands, were eligible for inclusion in the study. Healthy individuals also included boy scout patrol leaders, owners of hunting dogs and recreational runners. All were invited to participate if they pursued recreational activities in high-risk areas for tick bites, such as gardens, forests, grasslands and dunes [22]. Thus, the healthy individuals in this study represented a subgroup of healthy individuals, with a high risk of tick exposure. The healthy individuals were further subdivided into two groups. The first group consisted of healthy individuals who had received antibiotic treatment for LB-related symptoms in the past, as they had reported in the Lyme-specific questionnaire, and were referred to as treated healthy individuals. The second group comprised all other healthy individuals and these were referred to as untreated healthy individuals.

All study participants were asked to complete a Lyme-specific questionnaire. This questionnaire included questions on tick bites, the presence of EM, antibiotic treatment for LB and self-reported complaints at the moment of inclusion and during possible earlier episodes of LB. Information regarding the clinical symptoms, pleocytosis and intrathecal antibody production during active disease of the LNB patients was extracted from the hospital information system. Healthy individuals were recruited only if they reported no complaints at the time of inclusion in the study.

**Antibody detection in serum and serum–CSF pairs**

For the detection of *Borrelia*-specific antibodies in serum, the C6 enzyme-linked immunosorbent assay (ELISA) (ImmuneTech, Boston, MA, USA) was used [23]. Equivocal and positive C6 ELISA results were confirmed using the recomLine immunoglobulin (Ig)M and IgG
immunoblot test (Mikrogen GmbH, Neuried, Germany) [24]. Detection of intrathecally produced Borrelia-specific IgM and IgG antibodies was performed using the second-generation IDEIA LNB test (Oxoid, Hampshire, UK), which was adapted from the original publication by Hansen et al. [25]. Most importantly, the dilution of CSF was adjusted from 1:10 to 1:5, and various incubation times (of patient samples, conjugate and substrate) were shortened. The C6 ELISA and the IDEIA LNB tests were performed using a DS2-automated ELISA instrument (Dynex Technologies, Chantilly, VA, USA) and analysed with the DS-Matrix™ software (Dynex Technologies). The immunoblot results were recorded with an automated recomScan software (Mikrogen GmbH). All assays were performed according to the instructions of the respective manufacturers and were interpreted as described previously [20].

Isolation of peripheral blood mononuclear cells

Isolation of peripheral blood mononuclear cells (PBMCs) was performed from whole blood specimens which were collected in lithium heparin tubes. If isolation of PBMCs started within 8 h after venipuncture, 3 ml of fresh, pre-warmed (37°C) Roswell Park Memorial Institute (RPMI) medium (Life Technologies, Invitrogen, Bleiswijk, the Netherlands) was added to 5 ml of blood and, after gently mixing, transferred into a Leucosep tube (Oxford Immunotec Ltd, Abingdon, UK). PBMCs were separated through density gradient centrifugation (Hettich Rotanta 460 RS; rotor 5624) at room temperature for 15 min at 1000 g. If isolation of PBMCs was performed between 8 and 32 h after venipuncture, then a T-Cell Xtend (Oxford Immunotec Ltd) step was performed prior to the addition of 3 ml of RPMI medium and density gradient centrifugation, as previously described [20,26,27]. After centrifugation, the PBMC fraction was collected and washed twice in 10 ml RPMI medium. The first wash step was performed at room temperature for 7 min at 600 g; the second wash step was also performed at room temperature for 7 min at 300 g. If necessary, excess erythrocytes were removed between the first and second wash step using human erythrocyte lysis buffer [0.010 M KHCO3, 0.0001 M ethylenediamine tetraacetic acid (EDTA), 0.150 M NH4Cl (pH 7.3 ± 0.1)]. After addition of 5 ml of lysis buffer, the solution was incubated for 5 min at 2°C and subsequently centrifuged using the first wash step centrifugation program. The final pellet was suspended in 1.1 ml of fresh, prewarmed (37°C) AIM-V medium (Life Technologies) and PBMCs were counted using the AC.T diff 2 analyser (Beckman Coulter, Woerden, the Netherlands), as described previously [20]. After isolation, the PBMCs were adjusted to a concentration of 2.5 × 10⁶/ml using AIM-V medium, 100 µl of which (2.5 × 10⁵ PBMCs) was tested in the in-house Borrelia ELISpot assay and the commercial LymeSpot assay [Autoimmun Diagnostika (AID) GmbH, Straßberg, Germany].

The in-house Borrelia ELISpot assay

The in-house Borrelia ELISpot assay was performed as previously described [20]. In brief, a precoated polyvinylidene difluoride (PVDF) ELISpotPro 96-well plate (Mabtech, Nacka Strand, Sweden) was used, and four wells were tested for each study participant. These wells contained 50 µl of positive control [anti-human CD3 monoclonal antibody (mAb) CD3-2 (0.1 µg/ml); Mabtech], 50 µl of negative control (AIM-V medium), 50 µl of B. burgdorferi B31 whole cell lysate (5 µg/ml; AID), hereafter referred to as B. burgdorferi B31, and 50 µl of outer surface protein (Osp)-mix (5 µg/ml; AID), respectively, which were used to stimulate the PBMCs. The Osp-mix consisted of a pool of 9-mer to 11-mer peptides of Osp-A (B. burgdorferi, B. afzelii and B. garinii), native Osp-C (B. afzelii) and recombinant p18. For the current study, this protocol was extended by the addition of two wells: the first additional well contained 100 µl of B. burgdorferi B31 (5 µg/ml) and the second additional well contained 100 µl of Osp-mix (5 µg/ml) to stimulate the PBMCs (Supporting information, Table S1).

The numbers of Borrelia-specific interferon (IFN)-γ-secreting T cells/2.5 × 10⁵ PBMCs (displayed as black spots) were measured with an ELISpot reader (AID) and counted by two different people using the ELISpot software (AID), hereafter referred to as the numbers of spot-forming cells (SFCs). SFCs were counted without prior knowledge of the medical background of the study participants. The SFC size used was based on the expected SFC size of an IFN-γ-producing T cell, as determined by Feske et al. [28], and was set on –2.8 log (mm²). Samples that had a discrepancy in the numbers of SFCs between the two counting persons were recounted by a third person, whose result was leading. For samples that were stimulated with 50 µl of Borrelia antigen, the conditions for recounting have been described previously [20]. For samples which were stimulated with 100 µl of B. burgdorferi B31, a recount was performed for those samples which had a discrepancy in the numbers of SFCs in the critical area (between 0 and 10 SFCs), determined by receiver operating characteristic (ROC) curve analysis. When 100 µl of Osp-mix was used, those samples which had a discrepancy in the numbers of SFCs in the critical area (between 0 and 5 SFCs), determined by ROC curve analysis, were recounted. The results of the in-house Borrelia ELISpot assay were only interpreted when the assay was valid; i.e. when the numbers of SFCs upon stimulation in the positive control well were ≥ 20 and in the negative control well were ≤ 6 (the latter representing spontaneous SFCs) (Supporting information, Table S1). If the assay was valid,
the final numbers of SFCs in the *Borrelia* antigen-stimulated wells were determined. For the wells containing 50 µl of *Borrelia* antigen, this was performed by subtraction of the numbers of SFCs in the negative control well from the numbers of SFCs in the *Borrelia* antigen-stimulated well. For the wells containing 100 µl of *Borrelia* antigen, the final numbers of SFCs were calculated by first multiplying the numbers of SFCs in the negative control well by 2 before subtracting them from the numbers of SFCs in the *Borrelia* antigen-stimulated well (Supporting information, Table S1). The final numbers of SFCs corresponded with the numbers of SFCs after stimulation with either *B. burgdorferi* B31 or Osp-mix. For some cases, the *Borrelia* antigens were tested several times and, for such cases, the median T-cell count was used to determine the final numbers of SFCs. Using the extended version of our in-house *Borrelia* ELISpot assay, we were able to compare our in-house *Borrelia* ELISpot assay with the LymeSpot assay on the basis of exactly the same (absolute) amount of *Borrelia* antigens (100 µl of a 5 µg/ml concentration per well), as prescribed in the LymeSpot assay protocol. In addition, we could also study the effect of various amounts of *Borrelia* antigen (50 versus 100 µl of a concentration of 5 µg/ml) on the numbers of SFCs for the in-house *Borrelia* ELISpot assay (Supporting information, Table S1).

**The LymeSpot assay**

The LymeSpot assay (AID) was run in parallel with the in-house *Borrelia* ELISpot assay. The LymeSpot assay uses a 96-well PVDF plate coated with anti-human IFN-γ antibodies. The assay was performed according to the manufacturer’s protocol (Supporting information, Table S1), except for the isolation of the PBMCs and the amount of PBMCs/well, for which our standard protocol was used as described above and in Supporting information, Table S1. In a pilot study we investigated the influence of this deviation from the LymeSpot protocol, and showed that this had no impact on the diagnostic performance of the LymeSpot assay (see Supporting information, Data S4). Stimulation of the PBMCs in the LymeSpot assay was performed using a negative control (100 µl of AIM-V medium), a positive control (100 µl of Pokeweed; AID), 100 µl of *B. burgdorferi* B31 (5 µg/ml; AID) and 100 µl of Osp-mix (5 µg/ml; AID). Both the *B. burgdorferi* B31 and the Osp-mix antigens were identical to the *Borrelia* antigens used for the in-house *Borrelia* ELISpot assay described above. If the PBMC yield was sufficient, both controls and antigens were tested in duplicate (Supporting information, Table S1).

The final LymeSpot results were only calculated when the assay was valid. Following the manufacturer’s instructions, the LymeSpot results were valid when the positive control well had ≥ 50 SFCs and the negative control well had ≤ 10 SFCs. The final LymeSpot results were calculated in two ways. First, the average numbers of SFCs were calculated, similarly as described above for the in-house *Borrelia* ELISpot assay, to allow an objective comparison of the results of the LymeSpot assay with those of the in-house *Borrelia* ELISpot assay (Supporting information, Table S1). Secondly, stimulation indices (SIs) were calculated following the protocol of the manufacturer (Supporting information, Fig. S2). For this, the numbers of SFCs of the negative control needed to be established first. If these numbers were between 3 and 10, SIs were calculated by dividing the numbers of *Borrelia*-specific SFCs by the numbers of SFCs of the negative control. If the numbers of SFCs of the negative control were between 0 and 2, SIs were calculated by dividing the final numbers of *Borrelia*-specific SFCs by 1. The final LymeSpot results were based on the combination of the results of the SIs of both the *B. burgdorferi* B31 and the Osp-mix antigens, and a case could either be categorized as negative, positive (highly specific), or require diagnostic verification (Supporting information, Fig. S2).

**Data handling and statistical analysis**

The results of the in-house *Borrelia* ELISpot assay using 50 µl and using 100 µl of *Borrelia* antigen and the results of the LymeSpot assay were compared with regard to their ability to detect active LNB patients and to distinguish them from the study participants in the other three groups. The 50-µl results were published previously, as part of a larger study population (n = 243) [20]. For both ELISpot assays, a comparison was performed based on the individual, as well as the combined results of the numbers of SFCs after stimulation with either 50 or 100 µl *B. burgdorferi* B31 and 50 or 100 µl Osp-mix. In addition, for the LymeSpot assay, the *B. burgdorferi* B31-specific SI, the Osp-mix-specific SI and the final results based on the combination of both SIs (Supporting information, Fig. S2) were compared between the four study groups.

Dichotomous, unrelated data were analysed using the χ² or Fisher’s exact test. Quantitative, unrelated data comparing more than two groups were analysed using the Kruskal–Wallis χ² test, and subsequent two-group comparisons were analysed using the Dunn’s test [29]. Quantitative, unrelated data comparing two groups were analysed using the Wilcoxon rank sum test. Quantitative, related data comparing greater than or equal to two groups were analysed using the Wilcoxon signed-rank test with continuity correction.

To assess the diagnostic performance of both ELISpot assays, various ROC curves were constructed and used to calculate the area under the curve (AUC), sensitivity, specificity, positive predictive value (PPV) and negative predictive value (NPV) with 95% confidence intervals (CIs) [30]. The optimal threshold was calculated using
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the point on each ROC curve for which the distance to the upper left corner (where both sensitivity and specificity are 100%) was shortest, and was determined by the square root of [(1-sensitivity)²+(1-specificity)²]. The sensitivity, specificity, PPV and NPV were calculated based on the optimal threshold. For both ELISPOT assays, a ROC curve was constructed for each antigen separately by comparing the numbers of Borrelia-specific SFCs among active LNB patients with those among the other three groups, as well as for the results of both antigens together. To assess the diagnostic performance of each ELISPOT assay using the results of both antigens together, a binomial logistic regression (BLR) model was built before the ROC curves were constructed. The numbers of B. burgdorferi B31-specific SFCs and the numbers of Osp-mix-specific SFCs, without and with their interaction term, were included as predictor variables in the model; the outcome variable was binary: sick (all active LNB patients) or not-sick (all other study participants). The performance of the BLR model was assessed by calculating the prediction error using cross-validation. For the LymeSpot assay, ROC curve analysis was also performed based on the SIs after stimulation with B. burgdorferi B31 and Osp-mix, as described in the Materials and methods section covering the LymeSpot assay and in Supporting information, Fig. S2. The final results that needed diagnostic verification were classified as ‘positive’, and were combined with the positive results. Comparison of the ROC curves was performed using DeLong’s test for two correlated ROC curves [30].

Raw P-values < 0.05 were interpreted as statistically significant, which were subsequently followed by two-group comparisons where appropriate. To account for the multiple statistical analyses in this study, we applied the Benjamini–Hochberg (BH) procedure and controlled the false discovery rate at the level of 2.5%, i.e. no more than one false positive was allowed to be found in our list of rejections [31]. For all statistical analyses and construction of the figures, Rstudio version 1.1.383, 2009-2017 (Rstudio, Boston, MA, USA) was used.

Results

Study population

Ninety-two study participants were eligible for inclusion in the study; however, five (5.4%) patients were excluded. Four study participants, one active LNB patient, one untreated and two treated healthy individuals, were excluded because of insufficient amounts of PBMCs to perform both ELISPOT assays. One untreated healthy individual was excluded because the negative control well in the in-house Borrelia ELISPOT assay was invalid (> 6 SFCs). Eighty-four (96.6%) of the 87 study participants comprised a subgroup of our recently published study [20]; 77 (88.5%) of the 87 study participants were also part of another published study [21] (Supporting information, Table S3).

Characteristics of the active LNB patients

Eighteen active LNB patients were included in the study, with a median of 6.0 days after the start of antibiotic treatment for their active disease; their median age was 54.7 years (Table 1). Based on the EFNS criteria [10], 12 of 18 (66.7%) active LNB patients were classified as definite LNB cases and the remaining six (33.3%) were classified as possible LNB cases because of the lack of intrathecally produced Borrelia-specific antibodies (Table 1). Clinical symptoms consisted of radiculopathy (n = 2), cranial neuropathy (n = 7) or central nervous system disease (n = 5). Four patients had a combination of different symptoms: one patient had radiculopathy and cranial neuropathy, one patient had radiculopathy, cranial and peripheral neuropathy, one patient had radiculopathy and central nervous system disease and the last patient had cranial neuropathy and central nervous system disease (data not shown). Most active LNB patients had Borrelia-specific antibodies in their blood [15 of 18 (83.3%)], which was greater compared to treated LNB patients [one of 12 (8.3%)] and untreated healthy individuals [seven of 47 (14.9%)] (adjusted P-value ≤ 0.002) (Table 1). The majority [13 of 18 (72.2%)] of the active LNB patients showed complete recovery after the end of antibiotic therapy for active LNB, which was assessed by the neurologist with a median of 38.0 days after antibiotic treatment ended (Table 1).

Characteristics of the treated LNB patients

Twelve treated LNB patients were included in the study, who were diagnosed with active LNB on average 5.4 years ago (Table 1). The median age of the treated LNB patients at inclusion was 56.3 years and the majority (91.7%) were classified as definite LNB cases at the time of diagnosis of active LNB in the past. One (8.3%) patient was classified as a possible LNB case because of the absence of pleocytosis (Table 1). Clinical symptoms included radiculopathy (n = 1), cranial neuropathy (n = 4) or central nervous system disease (n = 3). Four treated LNB patients had combined symptomatology: one patient had radiculopathy, cranial neuropathy and central nervous system disease, one patient had radiculopathy and cranial neuropathy, one patient had radiculopathy and peripheral neuropathy and one patient had cranial and peripheral neuropathy (data not shown). Ten (83.3%) of the 12 treated LNB patients showed complete recovery after the end of antibiotic therapy for active LNB in the past, which was assessed by the neurologist with a median of 37.0 days after antibiotic treatment ended (Table 1). At inclusion in this study, however, eight (66.7%) of the 12
treated LNB patients reported complaints in the Lyme-specific questionnaire (Table 1). These self-reported symptoms included fatigue, neuropathic complaints, myalgias, arthralgias and cognitive complaints (data not shown).

**Characteristics of the healthy individuals**

A total of 57 healthy individuals were included. Ten (17.5%) reported having had antibiotic treatment for an early manifestation of LB in the past, which took place on average 5.0 years ago, and who were therefore classified as treated healthy individuals (Table 1). The median age of the treated healthy individuals was 55.2 years. Nine (90.0%) of the 10 treated healthy individuals reported having had a tick bite, and although this percentage was higher than among the other three groups, it was not statistically significant. Eight (80.0%) of the treated healthy individuals reported an EM, which was higher than among active LNB patients [one of 18 (5.6%)] and untreated healthy individuals [two of 47 (4.3%)] (adjusted P-value ≤ 0.002) (Table 1). The other two either reported flu-like symptoms or an atypical skin rash after the tick bite.

The remaining 47 (82.5%) healthy individuals all reported never to have had antibiotic treatment for LB, and thus
Table 2. Comparison of the ELISpot results expressed in the numbers of spot-forming cells for the in-house *Borrelia* ELISpot assay and the LymeSpot assay among the four study groups

| Study participants | *Borrelia* antigen | SFC count (median; IQR) | Statistics |
|--------------------|-------------------|-------------------------|------------|
|                    |                   | 50 µl | 100 µl | 100 µl | 100 µl | 100 µl | 100 µl |
|                    |                   | In-house *Borrelia* ELISpot assay | LymeSpot assay | 50 µl in-house *Borrelia* ELISpot assay versus 100 µl in-house *Borrelia* ELISpot assay | 100 µl in-house *Borrelia* ELISpot assay versus 100 µl LymeSpot assay |
| All study participants | Bb B31 | 2.0 (0.8–6.0) | 4.0 (2.0–9.5) | 5.0 (2.0–10.3) | < 0.001 | 0.006<sup>b</sup> | 0.360 | > 0.025 |
| (n = 87) | Osp-mix | 1.0 (0.0–2.0) | 1.0 (0.0–3.0) | 1.5 (0.5–3.3) | 0.786 | > 0.025 | 0.685 | > 0.025 |
| Active LNB patients | Bb B31 | 5.5 (1.3–7.8) | 6.5 (1.5–11.5) | 6.1 (2.6–10.2) | > 0.025 | > 0.025 | > 0.025 | > 0.025 |
| (n = 18) | Osp-mix | 0.0 (0.0–4.8) | 1.0 (0.0–3.0) | 1.3 (0.1–2.5) | 0.017<sup>c</sup> | ≤ 0.005<sup>d</sup> | > 0.025 | > 0.025 |
| Treated LNB patients | Bb B31 | 5.5 (2.0–9.9) | 9.5 (5.0–17.5) | 8.4 (2.4–12.8) | > 0.025 | > 0.025 | > 0.025 | > 0.025 |
| (n = 12) | Osp-mix | 0.5 (0.0–4.5) | 0.5 (0.0–2.5) | 1.5 (0.9–3.3) | 0.018 | 0.017<sup>e</sup> | 0.014<sup>e,f</sup> | > 0.025 |
| Treated healthy individuals (n = 10) | Bb B31 | 9.3 (3.5–27.0) | 6.5 (3.3–22.2) | 26.1 (7.5–73.7) | > 0.025 | > 0.025 | > 0.025 | > 0.025 |
| Untreated healthy individuals (n = 47) | Osp-mix | 2.0 (1.1–6.0) | 2.5 (0.5–7.8) | 5.6 (3.3–12.0) | > 0.025 | > 0.025 | > 0.025 | > 0.025 |
| Bb B31 versus Osp-mix | BH<sup>a</sup> (overall) | < 0.001 | < 0.001 | < 0.001 | > 0.025 | 0.017<sup>c</sup> | ≤ 0.005<sup>d</sup> | > 0.025 |
| Bb B31 | BH<sup>a</sup> (within each group) | > 0.025 | 0.017<sup>c</sup> | ≤ 0.005<sup>d</sup> | > 0.025 | > 0.025 |
| Osp-mix | BH (overall) | 0.018 | 0.132 | 0.075 | > 0.025 | > 0.025 |
| Bb B31 | BH (two-group) | 0.015<sup>e</sup> | > 0.025<sup>f</sup> | 0.014<sup>e,f</sup> | > 0.025 | > 0.025 |
| Osp-mix | BH (overall) | 0.489 | 0.766 | 0.058 | > 0.025 | > 0.025 |
| Bb B31 | BH (two-group) | n.d.<sup>g</sup> | n.d.<sup>g</sup> | 0.005<sup>e,f</sup> | > 0.025 | > 0.025 |

*SFC* = spot-forming cell; IQR = interquartile range; ELISpot = enzyme-linked immunosorbent spot; BH = Benjamini–Hochberg; n = number of study participants; Bb B31 = *Borrelia burgdorferi* B31; Osp = outer surface protein; n.d. = not done; LNB = Lyme neuroborreliosis.

<sup>a</sup>To correct for multiple comparisons, the Benjamini-Hochberg procedure was applied with a false discovery rate of 2.5%; <sup>b</sup>the numbers of SFCs among untreated healthy individuals were significantly higher after stimulation with 100 µl of *B. burgdorferi* B31 compared to 50 µl of *B. burgdorferi* B31; <sup>c</sup>the numbers of SFCs after stimulation with 100 µl of *B. burgdorferi* B31 among active Lyme neuroborreliosis (LNB) patients were significantly higher compared to stimulation with 100 µl of Osp-mix; <sup>d</sup>the numbers of SFCs after stimulation with 100 µl of *B. burgdorferi* B31 among untreated healthy individuals and active LNB patients were significantly higher compared to stimulation with 100 µl of Osp-mix (adjusted P-values are <0.001 and 0.005, respectively); <sup>e</sup>untreated versus treated healthy individuals; <sup>f</sup>as the initial comparison was significantly different (raw P-value <0.050), two-group comparisons were also performed; <sup>g</sup>as the initial comparison was not significantly different (raw P-value >0.050), two-group comparisons were not performed.
were classified as untreated healthy individuals. Their median age was 35.1 years, which was younger than the other three groups (adjusted P-value ≤ 0.018) (Table 1).

**Influence of the different amounts of *Borrelia* antigen used on the median numbers of SFCs in the in-house *Borrelia* ELISpot assay**

PBMCs of all 87 study participants were stimulated with 50 µl [20] and 100 µl of *B. burgdorferi* B31 and Osp-mix, and subsequently tested in our in-house *Borrelia* ELISpot assay (Table 2). Overall, when 50 µl of *B. burgdorferi* B31 was used to stimulate the PBMCs, a lower median number of SFCs was obtained than when 100 µl of antigen was used (2.0 versus 4.0) (adjusted P-value < 0.001) (Table 2). When the four study groups were analysed separately, the association between the use of lower amounts of antigen as stimulant and the lower median number of SFCs remained for untreated healthy individuals (1.5 versus 2.0) (adjusted P-value = 0.006) (Table 2). Stimulation of PBMCs with either 50 or 100 µl of Osp-mix did not result in a difference between the median numbers of SFCs in the in-house *Borrelia* ELISpot assay (1.0 versus 1.0) (adjusted P-value = 0.786) (Table 2).

**Influence of the different *Borrelia* antigens used for PBMC stimulation on the median numbers of SFCs in the two ELISpot assays**

Analysis of the results of all 87 study participants showed that PBMC stimulation with 50 µl of *B. burgdorferi* B31 resulted in a higher median number of SFCs than stimulation with 50 µl of Osp-mix in the in-house *Borrelia* ELISpot assay (2.0 versus 1.0) (adjusted P-value < 0.001) (Table 2). These results are similar to the results we have published previously using a study population of 243 study participants [20]. When the four study groups were analysed separately, the median numbers of *B. burgdorferi* B31-specific SFC counts were higher compared to the median numbers of Osp-mix-specific SFC counts, although not significant (adjusted P-values > 0.025) (Table 2). A higher median number of SFCs after PBMC stimulation with *B. burgdorferi* B31 compared to PBMC stimulation with Osp-mix was also seen when a volume of 100 µl of *Borrelia* antigen was used in the in-house *Borrelia* ELISpot assay (4.0 versus 1.0) (adjusted P-value < 0.001) (Table 2). Comparisons within each of the four groups showed that this difference remained significant for active LNB patients (adjusted P-value = 0.017) (Table 2).

In the LymeSpot assay, the higher yield of the *B. burgdorferi* B31 over the Osp-mix remained when the median numbers of SFCs were compared (5.0 versus 1.5) (adjusted P-value < 0.001) (Table 2). When the four study groups were analysed separately, *B. burgdorferi* B31 remained superior in the LymeSpot assay for untreated healthy individuals and active LNB patients (adjusted P-values ≤ 0.005) (Table 2).

**Comparison of the median numbers of SFCs in the two ELISpot assays between the four study groups after stimulation of the PBMCs with *B. burgdorferi* B31**

The PBMCs of treated healthy individuals were stimulated the most when either 50 µl of *B. burgdorferi* B31 was used in the in-house *Borrelia* ELISpot assay or 100 µl of *B. burgdorferi* B31 was used in the LymeSpot assay. The PBMCs of treated LNB patients were stimulated the most when 100 µl of *B. burgdorferi* B31 was used in the in-house *Borrelia* ELISpot assay (Table 2; Fig. 1a,c,e). In contrast, the PBMCs of untreated healthy individuals were stimulated the least, irrespective of the volume and the ELISpot assay used. An increased T-cell activation for patients and treated healthy individuals after PBMC stimulation with *B. burgdorferi* B31 was also seen in our previous study, which included more study participants (n = 243), and suggests that the ELISpot activity is related to exposure to the *Borrelia* bacterium [20].

Analysis of the different amounts of *B. burgdorferi* B31 showed that when 50 µl was used to stimulate the PBMCs in the in-house *Borrelia* ELISpot assay, the median number of SFCs of 1.5 for untreated healthy individuals was lower compared to the median number of SFCs of 9.3 for treated healthy individuals (adjusted P-value = 0.015) (Table 2, Fig. 1a). When 100 µl of *B. burgdorferi* B31 was used to stimulate the PBMCs, no differences were found between the four study groups for the in-house *Borrelia* ELISpot assay (adjusted P-value > 0.025) (Table 2, Fig. 1c). For the LymeSpot assay, the results were only significantly different between untreated (less ELISpot activity) and treated healthy individuals (more ELISpot activity) (adjusted P-value = 0.014) (Table 2, Fig. 1e).

Overall, no difference was found between the median numbers of SFCs between both ELISpot assays when 100 µl of *B. burgdorferi* B31 was used to stimulate the PBMCs (adjusted P-value = 0.360). Similarly, no differences were found when the four study groups were analysed separately (adjusted P-values > 0.025) (Table 2).

**Comparison of the median numbers of SFCs in the two ELISpot assays between the four study groups after stimulation of the PBMCs with Osp-mix**

Similar to stimulation with *B. burgdorferi* B31, the PBMCs of treated healthy individuals were activated most upon stimulation with Osp-mix (Table 2, Fig. 1b,d,f). No significant differences between the four study groups were found for the in-house *Borrelia* ELISpot assay using 50 or 100 µl of Osp-mix (adjusted P-value 0.489 and 0.766, respectively) (Table 2, Fig. 1b,d). For the LymeSpot assay, however, stimulation of the PBMCs with 100 µl of
Osp-mix resulted in a significantly higher median number of SFCs of 5.6 for treated healthy individuals compared to the median number of SFCs of 1.0 for untreated healthy individuals (adjusted $P$-value = 0.005) (Table 2, Fig. 1f).

Similar to the use of 100 µl of $B. burgdorferi$ B31, no difference was seen between the median numbers of SFCs between both ELISpot assays upon stimulation of the PBMCs with 100 µl of Osp-mix (adjusted $P$-value = 0.685).

Fig. 1. Results of the in-house $Borrelia$ enzym-linked immunosorbent spot (ELISpot) assay (a–d) and the LymeSpot assay (e–f) expressed in the numbers of spot-forming cells (SFCs). (a) (50 µl), (c) and (e) (both 100 µl) are the results after peripheral blood mononuclear cell (PBMC) stimulation with $Borrelia burgdorferi$ B31, and (b) (50 µl), (d) and (f) (both 100 µl) are the results after PBMC stimulation with outer surface protein (Osp)-mix among active Lyme neuroborreliosis patients (ANB), treated Lyme neuroborreliosis patients (TNB), treated healthy individuals (THI) and untreated healthy individuals (UHI). The displayed $P$-values are corrected and interpreted using the Benjamini–Hochberg procedure with a false discovery rate of 2.5% for multiple comparisons (only false discovery rates < 0.025 are displayed).
Subsequent comparisons within each group also did not show a difference (adjusted $P$-values $>$ 0.025) (Table 2).

The diagnostic performance of the two ELISpot assays based on the numbers of SFCs

The diagnostic performance of the in-house *Borrelia* ELISpot assay and the LymeSpot assay were evaluated using ROC curve analysis, for which the numbers of SFCs were used. In order to enable a fair comparison between the two assays, the results obtained with PBMCs that were stimulated with 100 µl of *Borrelia* antigen were used for the in-house *Borrelia* ELISpot assay and compared with the results of the LymeSpot assay. The results obtained with 100 µl of *Borrelia* antigen were used, as this is the standard in the LymeSpot assay (Supporting information, Table S1). ROC curves were constructed based on the results obtained after PBMC stimulation with the *B. burgdorferi* B31 and the Osp-mix separately, as well as on the combined results of both *Borrelia* antigens. The calculated AUCs based on the individual *Borrelia* antigens were comparable to a random predictor, and ranged from 0.459 to 0.570 (Table 3, Fig. 2a,b). No difference was found between the AUC of the in-house *Borrelia* ELISpot assay and the AUC of the LymeSpot assay based on the numbers of *B. burgdorferi* B31-specific SFCs (AUC = 0.553 and 0.570, respectively) (adjusted $P$-value = 0.974) (Table 3, Fig. 2a). Similarly, comparison of the AUCs from the two ELISpot assays based on the numbers of Osp-mix-specific SFCs also showed no difference (AUC = 0.479 for the in-house *Borrelia* ELISpot assay and AUC = 0.459 for the LymeSpot assay, respectively) (adjusted $P$-value = 0.930) (Table 3, Fig. 2b).

Calculation of the optimal thresholds for the two assays using a single *Borrelia* antigen showed that the sensitivity and NPV was highest for the LymeSpot assay when *B. burgdorferi* B31 was used to stimulate the PBMCs (sensitivity = 66.7%, NPV = 87.0%) (Table 3). The specificity was highest for the in-house *Borrelia* ELISpot assay irrespective of whether *B. burgdorferi* B31 or Osp-mix was used to stimulate the PBMCs (66.7% each) (Table 3). The PPV was highest for the in-house *Borrelia* ELISpot assay when *B. burgdorferi* B31 was used to stimulate the PBMCs (60.6%) (Table 3).

Combining the results of the two *Borrelia* antigens without or with their interaction term as risk factors in a BLR model also resulted in AUCs that were comparable to a random predictor (range = 0.429–0.549) (Table 3). Comparison of the AUCs for the in-house *Borrelia* ELISpot assay and the LymeSpot assay without their interaction term did not show a difference (AUC = 0.546 and 0.429, respectively) (adjusted $P$-value = 0.517) (Table 3, Fig. 2c). Similarly, the AUCs for the in-house *Borrelia* ELISpot assay and the LymeSpot assay with their interaction term were also comparable (AUC = 0.549 and 0.521, respectively) (adjusted $P$-value = 0.959) (Table 3, Fig. 2c). For both ELISpot assays, the prediction errors of the BLR models with the interaction term were only slightly better than the prediction errors of the BLR models without the interaction term (20.7% for both versus 21.8% for both) (Table 3). Thus, approximately one in five patients were wrongly diagnosed by using the BLR models.

Furthermore, no differences were found between the AUCs of both BLR models for the in-house *Borrelia* ELISpot assay (adjusted $P$-value = 1.000) as well as for the LymeSpot assay (adjusted $P$-value = 0.717) (Table 3, Fig. 2c). The highest sensitivity was found for the in-house *Borrelia* ELISpot assay when both antigens without their interaction term were included in the BLR model (66.7%); the highest specificity (72.5%) and PPV (33.3%) was found for the in-house *Borrelia* ELISpot assay when both antigens with their interaction term were included in the BLR model. The NPV was highest for the in-house *Borrelia* ELISpot assay, irrespective of whether or not the interaction term was included (85.7% each) (Table 3).

In conclusion, the two ELISpot assays showed a poor diagnostic performance for the diagnosis of active LNB when the numbers of SFCs were used in the ROC curve analyses, with sensitivities ranging from 44.4 to 66.7%, specificities from 42.0 to 72.5%, PPVs from 21.8 to 33.3% and NPVs from 80.5 to 87.0% (Table 3).

The diagnostic performance of the LymeSpot assay based on the SIs

ROC curve analysis based on a combination of the SIs of both antigens following the manufacturer’s protocol (Supporting information, Fig. S2) resulted in an AUC of 0.487 (Table 3), which almost perfectly fitted the random predictor (Fig. 2d). Based on this analysis, the LymeSpot assay had a sensitivity of 27.8% to diagnose active LNB (Table 3). Thus, only five of the 18 active LNB patients were correctly identified using the LymeSpot assay (Table 4). Of these five active LNB patients, two had a positive LymeSpot result, and for three active LNB patients the LymeSpot results would still need diagnostic verification according to the manufacturer’s instructions. The remaining 13 (72.2%) active LNB patients had a negative LymeSpot result (Table 4). When the results that needed diagnostic verification were excluded from the positive results, the sensitivity of the LymeSpot assay decreased to 11.1%. The specificity of the LymeSpot assay was 69.9% (Table 3), and 21 (30.4%) of the 69 controls either needed diagnostic verification $[n = 11 (15.9%)]$ or had a positive LymeSpot result $[n = 10 (14.5%)]$ (Table 4). Interestingly, the percentage of positive LymeSpot results was highest among treated healthy individuals $[seven of 10 (70.0%)]$; however, this was not significantly higher when compared to the other groups (adjusted $P$-values $>$ 0.025) (Table 4).
Table 3. Diagnostic performance of the in-house *Borrelia* ELISpot assay and the LymeSpot assay based on the numbers of spot-forming cells obtained after peripheral blood mononuclear cell stimulation with *Borrelia burgdorferi* B31 and Osp-mix separately, and based on the combined numbers of spot-forming cells of both *Borrelia* antigens by using a binomial logistic regression model.

| Interpretation | ELISpot assay | Threshold \(a\) (95% CI) | AUC \(b\) (95% CI) | Sensitivity \(c\) (95% CI) | Specificity \(d\) (95% CI) | PPV \(e\) (95% CI) | NPV \(f\) (95% CI) | BH \(g\) Prediction error BLR model (%) |
|----------------|---------------|--------------------------|------------------|------------------|------------------|----------------|----------------|---------------------------------|
| SFC count after stimulation with Bb B31 (100 µl) | In-house *Borrelia* ELISpot assay | 7.0 (2.5-8.5) | 0.553 (0.382-0.714) | 61.1 (38.9-77.8) | 66.7 (42.0-81.2) | 30.6 (20.5-44.8) | 85.7 (79.1-92.2) | 0.974 |
| | LymeSpot assay | 5.3 (5.0-8.4) | 0.570 (0.433-0.706) | 66.7 (44.4-88.9) | 59.4 (44.9-72.5) | 29.7 (21.1-39.5) | 87.0 (79.6-94.7) | |
| SFC count after stimulation with Osp-mix (100 µl) | In-house *Borrelia* ELISpot assay | 2.5 (0.5-5.5) | 0.479 (0.326-0.631) | 44.4 (16.7-72.2) | 66.7 (33.3-87.0) | 23.3 (13.8-40.0) | 80.9 (72.2-87.1) | 0.930 |
| | LymeSpot assay | 2.3 (0.3-4.3) | 0.459 (0.305-0.614) | 50.0 (22.2-77.8) | 56.5 (24.6-84.1) | 22.2 (14.6-36.4) | 80.5 (72.3-87.5) | |
| SFC count of both antigens without interaction term in a BLR model\(c,d\) | In-house *Borrelia* ELISpot assay | 0.21 (0.20-0.22) | 0.546 (0.398-0.694) | 66.7 (38.9-88.9) | 49.3 (33.3-81.2) | 26.5 (20.0-40.0) | 85.7 (78.4-94.3) | 21.8 |
| | LymeSpot assay | 0.21 (0.21-0.22) | 0.429 (0.292-0.566) | 61.1 (27.8-88.9) | 42.0 (20.3-73.9) | 21.8 (16.1-29.8) | 80.8 (72.7-90.6) | 0.517 |
| SFC count of both antigens with interaction term in a BLR model\(c,d\) | In-house *Borrelia* ELISpot assay | 0.21 (0.19-0.22) | 0.549 (0.380-0.719) | 55.6 (33.3-77.8) | 72.5 (42.0-85.5) | 33.3 (21.1-50.0) | 85.7 (79.6-91.8) | 20.7 |
| | LymeSpot assay | 0.21 (0.19-0.22) | 0.482 (0.372-0.670) | 50.0 (33.3-83.3) | 68.1 (30.4-81.2) | 27.6 (18.9-41.7) | 83.9 (77.4-91.3) | 0.959 |
| Final result based on the stimulation indices of both antigens | LymeSpot assay | n.a. \(e\) | 0.487 (0.367-0.606) | 27.8 (11.1-50.0) | 69.6 (58.0-79.7) | 19.1 (7.7-32.0) | 78.6 (73.7-84.0) | n.a. |

ELISpot = enzyme-linked immunosorbent spot; AUC = area under the curve; PPV = positive predictive value; NPV = negative predictive value; CI = confidence interval; BH = Benjamini-Hochberg; BLR = binomial logistic regression; SFC = spot-forming cell; Bb B31 = *B. burgdorferi* B31; Osp = outer surface protein; n.a. = not applicable.

\(a\) The threshold is based on the numbers of spot-forming cells when the result of a single *Borrelia* antigen was analysed, and on the linear predictors of the binary logistic regression model when the combined results of both *Borrelia* antigens were analysed;\(b\) to correct for multiple comparisons, the Benjamini–Hochberg procedure was applied with a false discovery rate of 2.5%;\(c\) no significant difference (adjusted \(P\)-value = 1.000) between the outcome of the BLR model without interaction term and the outcome of the BLR model with interaction term for the in-house *Borrelia* ELISpot assay;\(d\) no significant difference (adjusted \(P\)-value = 0.717) between the outcome of the BLR model without interaction term and the outcome of the BLR model with interaction term for the LymeSpot assay; \(e\) see Supporting information, Fig. S2.
In total, 26 (29.9%) of the 87 study participants reported symptoms at inclusion in this study; all active LNB patients ($n = 18$) and eight (66.7%) of the 12 treated LNB patients. Overall, no association was found between the presence of symptoms and the results of the various ELISpot protocols (Table 5a). For treated LNB patients, the number of complaints reported at the start of the study was also not associated with the ELISpot results, irrespective of the ELISpot protocol used (Table 5a). As the treated healthy individuals were only included when they did not report any symptoms at the start of the study, elevated SFC counts in this group could not be linked to symptoms. Similarly, 16 (28.1%) of the 57 healthy individuals had a
Table 4. Interpretation of the LymeSpot assay based on the stimulation indices according to the protocol of the manufacturer

| Study groups                  | NEG (n; %) | POS (n; %) | BH (overall) | BH (two-group) |
|-------------------------------|------------|------------|--------------|----------------|
| Active LNB patients (n = 18)  | 13 (72.2)  | 5 (27.8)   |              |                |
| Treated LNB patients (n = 12) | 7 (58.3)   | 5 (41.7)   | 0.066        | > 0.025*       |
| Treated healthy individuals (n = 10) | 3 (30.0) | 7 (70.0) |              |                |
| Untreated healthy individuals (n = 47) | 38 (80.9) | 9 (19.1) |              |                |
| Control group (n = 69) | 48 (69.6) | 21 (30.4) | n.a.         | n.a.           |

NEG = negative; POS = positive; n = number of study participants; BH = Benjamini–Hochberg; LNB = Lyme neuroborreliosis; n.a. = not applicable.

*The positive results include those results that needed diagnostic verification; †three (16.7%) of 18 active Lyme neuroborreliosis (LNB) patients required diagnostic verification; ‡three (25.0%) of 12 treated LNB patients required diagnostic verification; †The presence of *Borrelia*-specific IgM or IgG also was not associated with elevated SFC counts (adjusted *P*-values ranged from 0.199 to 1.000; data not shown).

Among active LNB patients, no significant association was found between the duration of symptoms prior to the blood sampling and the ELISpot results, irrespective of the ELISpot protocol used (Table 5b). For most active LNB patients, the antibiotic treatment had already started at the time of blood sampling; however, no association was found between the duration of antibiotic therapy prior to blood sampling and the ELISpot results using the various ELISpot protocols (Table 5c). Similarly, no association was found between the degree of recovery and the T-cell response of active LNB patients (Table 5d). For treated LNB patients, the degree of recovery was assessed at a median of 37.0 days after the end of antibiotic treatment for active disease in the past (approximately 5.4 years ago; Table 1), therefore, we did not compare the degree of recovery with the various ELISpot results obtained in the current study.

ELISpot results versus *Borrelia*-specific antibodies

In our previous study, elevated numbers of *B. burgdorferi* B31-specific T cells were significantly associated with the presence of *Borrelia*-specific serum antibodies [20]. In this study, which included a smaller number of study participants, comparison of the *B. burgdorferi* B31-specific SFC counts with the serology results showed a trend towards a combined B- and T-cell response when all study participants were analysed together, irrespective of the ELISpot protocol used (Table 6a). Within-group comparisons also showed a (non-significant) trend towards a combined B- and T-cell response, except for treated LNB patients, who showed elevated *B. burgdorferi* B31-specific SFC counts in the absence of *Borrelia*-specific antibodies. This was, again, in line with the results found in our previous study [20].

The presence of *Borrelia*-specific IgM or IgG also was not associated with elevated SFC counts (adjusted *P*-values ranged from 0.199 to 1.000; data not shown).

Among active LNB patients, no significant association was found between the intrathecal production of *Borrelia*-specific antibodies and the T-cell response (Table 5b), which was similar to the results of our previous study in which 33 active LNB patients were included [20]. We also did not find a difference among the active LNB patients between negative and positive IgM AI results or between negative and positive IgG AI results when compared to the various SFC counts (adjusted *P*-values ranged from 0.131 to 1.000; data not shown).

For treated LNB patients, the presence of intrathecally produced *Borrelia*-specific antibodies was determined at the time of active disease in the past, therefore, we did not compare these results with the ELISpot results using the various ELISpot protocols, as these were performed on average 5-4 years (Table 1), at the time the treated LNB patients were included in this study.

Discussion

In the current study, the diagnostic performance of two ELISpot assays to diagnose active LNB were compared. The final study population consisted of 87 participants and comprised 18 active and 12 treated LNB patients, 10 healthy individuals who were treated for an early (mainly cutaneous) manifestation of LB in the past and 47 untreated healthy individuals. Both our in-house *Borrelia* ELISpot assay and the LymeSpot assay showed a poor diagnostic performance based on the numbers of SFCs with AUCs ranging from 0.429 to 0.570. The corresponding sensitivities, specificities, PPVs and NPVs ranged from 44.4 to
Osp = outer surface protein; SFC = spot-forming cell; ELISpot = enzyme-linked immunosorbent spot; n = number of study participants; IQR = interquartile range; BH = Benjamini–Hochberg; NEG = negative; POS = positive; LNB = Lyme neuroborreliosis.

*Symptoms are defined as the presence of symptoms at the start of the study. For (un)treated healthy individuals and treated Lyme neuroborreliosis (LNB) patients, the presence of symptoms was assessed by the completion of a Lyme-specific questionnaire; (un)treated healthy individuals were only included if they did not report any symptoms at the start of the study. For active LNB patients, the presence of symptoms was extracted from the hospital information system; b to correct for multiple comparisons, the Benjamini–Hochberg procedure was applied with a false discovery rate of 2.5%; cin total eight (66.7%) of the 12 treated LNB patients reported complaints at the start of the study. For all treated LNB patients, the presence of complaints was reported on average 5.4 years after the diagnosis of active disease in the past (Table 1); dSymptom duration is defined as the number of days the study participant experienced complaints prior to blood sampling; ePost AB-treatment time is defined as the number of days between the start of antibiotic AB) treatment and blood sampling (median days; IQR) f the degree of recovery (recovery status) was assessed after a median of 38.0 days after the end of antibiotic therapy for active disease (Table 1).
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66.7%, 42.0 to 72.5%, 21.8 to 33.3% and 80.5 to 87.0%, respectively. The diagnostic performance of the LymeSpot assay, using so-called SIs following the manufacturer's protocol, resulted in a comparably low AUC of 0.487, with a corresponding sensitivity of 27.8%, a specificity of 69.6%, a PPV of 19.1% and a NPV of 78.6%. Our study showed that the two ELISpot assays, irrespective of the protocol used, cannot be used to diagnose LNB or to monitor antibiotic treatment success.

The results of the 87 study participants of the in-house Borrelia ELISpot assay after stimulation of the PBMCs with 50 µl of B. burgdorferi B31 in the current study represent a subset of the results of the 243 study participants published previously [20]. The SFC counts between the four study groups of the subgroup in this study were comparable with the SFC counts between the four study groups of the entire study population. Both studies showed significantly higher numbers of SFCs after stimulation with B. burgdorferi B31 for treated healthy individuals compared to untreated healthy individuals. Active LNB patients and treated LNB patients also showed higher numbers of SFCs after stimulation with B. burgdorferi B31 compared to untreated healthy individuals, although not significant in the current study. This is most probably explained by the lower number of study participants per group in the current study. The association between the B- and T-cell response that was found in our previous study was also seen in the current study, although it was not significant, most probably due to the smaller study population. The overall conclusion, that elevated numbers of SFCs are associated with a previous contact with the Borrelia bacterium [20], however, was confirmed and could not be linked to symptomology nor to the degree of recovery or to antibiotic treatment. Elevated IFN-γ levels among asymptomatic individuals and previous LB patients have also been found by others [32–35].

Comparison of the in-house Borrelia ELISpot assay results after PBMC stimulation with either 50 or 100 µl of B. burgdorferi B31 showed similar results for three of the four study groups. However, among untreated healthy individuals, significantly higher numbers of SFCs were seen when 100 µl was used. This could be explained by the relatively higher number of untreated healthy individuals compared to the number of study participants in the other three groups.

Consistent with our previous study [20], we found that the use of Osp-mix as a T-cell stimulant resulted in very low numbers of SFCs, and cannot be used in its current composition to distinguish active LNB patients from the three control groups. Other studies also described a reduced performance of recombinant antigens compared to whole cell lysates [13,36]. This may, in part, be explained by the number of different antigens present: (a mixture of various) recombinant antigens contains far less antigens than a whole cell lysate. Alternatively, recombinant antigens are more specific, therefore limiting the possibility of cross-reactivity. It is known that Borrelia-specific antibodies show cross-reactivity with other diseases [37] and that the bacterium shows high sequence homology with bacteria such as Treponema or Leptospiro. Cross-reactivity could theoretically result in higher numbers of SFCs when a whole cell lysate of B. burgdorferi B31 is used in patients with an active or previous infection caused by bacteria such as Treponema or Leptospiro, or in healthy individuals who carry non-pathogenic Treponema or Leptospiro species. Previously, we have tested two patients with active leptospirosis in our in-house Borrelia ELISpot assay, and one of them had high numbers of SFCs after stimulation with a whole cell lysate of B. burgdorferi B31 [20].

Overall, the numbers of SFCs after stimulation with B. burgdorferi B31 were also relatively low. In our experience, as well as described by others – for tuberculosis or cytomegalovirus infections – the numbers of IFN-γ-secreting T cells among exposed or infected individuals measured in an ELISpot assay using comparable amounts of PBMCs, ranging from 2.0 × 10⁵ to 2.5 × 10⁵, are generally much higher [40–42]. The lack of T-cell activity among the active LNB patients could be explained by the choice of Borrelia antigens. In the Netherlands, LNB is mainly caused by B. garinii and B. bavariensis [43] and less frequently by B. burgdorferi sensu stricto. As we have discussed previously [20], we do not believe that the use of B. burgdorferi B31 whole cell lysate in the ELISpot assay resulted in the poor performance of both ELISpot assays, as B. burgdorferi, B. garinii and B. bavariensis are closely related and share many antigens. Von Baehr et al. [13] evaluated three Borrelia species and did not find any difference. Nordberg et al. [44] used B. garinii as a stimulating agent, CSF instead of blood and nitrocellulose-bottomed ELISpot plates instead of PVDF-bottomed plates, and also did not find higher numbers of activated T cells in their ELISpot assay. The Osp-mix we used contained antigens derived from an LNB-associated strain (B. garinii); however, the Osp-mix was inferior compared to the use of B. burgdorferi B31, as discussed in the previous paragraph. The lack of T-cell activity might also be explained by the inability of the human host to develop an adequate immune response against the Borrelia bacterium or the ability of the Borrelia bacterium to escape or suppress the immune system [45,46]. It could also be due to the disease manifestation that was studied, as already debated previously [20], as LNB implies a local infection of the brain. Testing blood might thus be less suitable, as the
Table 6. Overview of the B- and T-cell response among the various study groups

| Study group                        | AI      | T. van Gorkom et al. | n (%) | 50 µl Bb B31 | 100 µl Bb B31 | 50 µl Osp-mix | 100 µl Osp-mix | Final LymeSpot result |
|------------------------------------|---------|----------------------|-------|--------------|---------------|---------------|---------------|-----------------------|
|                                    |         |  |     | SFC count | SFC count | SFC count | SFC count | SFC count | Median (IQR) BH | Median (IQR) BH | Median (IQR) BH | Median (IQR) BH | Median (IQR) BH | Median (IQR) BH | Median (IQR) BH |
| (a) All study participants combined (n = 87) | NEG      | 60 (69.0) | 2.0 (0.4–5.3) | 3.0 (1.8–6.0) | 3.4 (1.4–8.6) | 1.0 (0.0–2.0) | 1.0 (0.0–3.0) | 43 | 17 | 0.986 |
|                                    | POS      | 27 (31.0) | 5.0 (1.5–8.8) | 8.0 (3.0–14.3) | 7.0 (3.8–19.4) | 0.0 (0.0–4.5) | 1.0 (0.0–4.0) | 18 | 9 | 0.986 |
| Active LNB patients (n = 18)       | NEG      | 3 (16.7) | 5.0 (2.5–19.5) | 5.0 (2.5–28.0) | 11.0 (6.3–30.2) | 1.0 (0.5–5.0) | 5.0 (2.5–6.0) | 1.0 | 0.0 | 43 | 17 | 0.986 |
|                                    | POS      | 15 (83.3) | 6.0 (1.5–7.5) | 8.0 (2.0–11.0) | 6.0 (2.8–9.3) | 0.0 (0.0–4.5) | 1.0 (0.0–2.5) | 12 | 3 | 0.430 |
| Treated LNB patients (n = 12)      | NEG      | 11 (91.7) | 6.0 (2.0–11.2) | 7.0 (5.0–19.0) | 8.3 (2.3–15.2) | 1.0 (0.0–5.0) | 0.0 | 0.0 | 43 | 17 | 0.986 |
|                                    | POS      | 1 (8.3) | 2.0 | 12.0 | 8.5 | 1.000 | 0.355 | 1.0 | 1.000 | 0.471 |
| Treated healthy individuals (n = 10) | NEG      | 6 (60.0) | 4.5 (1.5–10.9) | 3.5 (3.0–4.8) | 8.5 (2.4–15.8) | 1.8 (0.4–2.0) | 2.0 (0.5–2.8) | 3.8 (2.8–6.2) | 3.3 | 3 | 0.471 |
|                                    | POS      | 4 (40.0) | 24.0 (12.5–33.5) | 45.0 (19.2–72.2) | 78.8 (66.8–83.1) | 9.5 (5.5–14.2) | 11.0 (6.8–13.6) | 122 (10.0–12.6) | 33 | 7 | 0.471 |
| Untreated healthy individuals (n = 47) | NEG      | 40 (85.1) | 1.3 (0.0–3.0) | 2.0 (1.0–4.3) | 3.0 (1.2–6.8) | 1.0 (0.0–1.3) | 0.0 (0.0–1.0) | 1.0 | 0.0 | 814 |
|                                    | POS      | 7 (14.9) | 3.0 (0.8–5.5) | 60.0 (10.0–112) | 5.0 (3.5–10.0) | 0.377 | 0.0 (0.0–1.0) | 0.785 | 0.679 | 0.814 |
| (b) Active LNB patients (n = 18)   | NEG      | 6 (33.3) | 3.5 (2.3–19.0) | 4.0 (1.5–26.4) | 4.3 (2.6–8.9) | 2.0 (0.0–5.5) | 0.5 (0.0–1.8) | 0.890 | 0.3 (0.0–2.0) | 0.625 |
|                                    | POS      | 12 (66.7) | 6.0 (4.8–7.3) | 8.0 (2.3–10.5) | 6.6 (4.6–10.4) | 0.0 (0.0–2.8) | 1.0 (0.0–3.5) | 1.5 (0.5–2.9) | 4 | 2 | 1.000 |

Bb B31 = B. burgdorferi B31; OSP = outer surface protein; SFC = spot-forming cell; LymeSpot = enzyme-linked immunosorbent spot; AI = antibody index; n = number of study participants; IQR = interquartile range; BH = Benjamini–Hochberg; NEG = negative; POS = positive; LNB = Lyme neuroborreliosis; AB = antibiotic treatment for Lyme borreliosis.

aThe final LymeSpot result is based on a combination of the stimulation indices of both antigens following the protocol of the manufacturer (Supporting information, Fig. S2); bto correct for multiple comparisons, the Benjamini–Hochberg procedure was applied with a false discovery rate of 2.5%.
immune cells could have migrated towards the central and/or peripheral nervous system [47,48]. The testing of CSF, in combination with blood, may be more suitable [49]. Furthermore, IFN-γ may not be the best marker to diagnose active LNB. It would be interesting to investigate whether other cytokines and/or chemokines could improve the ELISpot assays tested in this study. Recently, the LymeSpot assay has been adapted by the manufacturer by adding the detection of interleukin (IL)-2. However, no data are available yet with regard to the diagnostic performance of this modified LymeSpot assay.

For the LymeSpot assay, the PBMC isolation procedure used in this study deviated from the manufacturer’s (AID) recommended protocol. These deviations from the LymeSpot protocol were made in order for the technician to be able to perform and process the ELISpot assays simultaneously, and to minimize the differences between the assays to allow for a more fair comparison. The PBMCs used in the LymeSpot assay were thus isolated according to the same protocol that was already in use in our laboratory for the in-house Borrelia ELISpot assay [20] and for the T-SPOT.TB test [20,26,50]. Consequently, the PBMC isolation differed at four points compared to the instruction manual of the LymeSpot assay.

First, the medium to dilute the blood prior to PBMC isolation differed, as RPMI medium was used instead of phosphate-buffered saline (PBS). Secondly, Leucosep tubes were used for the isolation of PBMCs, while the LymeSpot protocol advises to use standard tubes with a Ficoll gradient. As a consequence, the centrifugation steps of the isolation procedure were adjusted based on the instruction manual supplied with the Leucosep tubes. As the isolation of PBMCs is based on a gradient, we do not believe that the altered centrifugation time resulted in a different PBMC yield. An increased centrifugation speed could, potentially, result in a higher PBMC yield, but this should not influence the results of the LymeSpot assay, as the amount of PBMCs per well is standardized. This is confirmed by others [51,52], who showed that PBMCs isolated by Leucosep tubes performed equally well in the ELISpot assay compared to PBMCs isolated using the Ficoll-gradient technique.

Thirdly, the centrifugation steps that were used to wash the PBMCs and the number of times the PBMCs were washed differed from the LymeSpot protocol. However, in the literature, various centrifugation speeds and times for washing the PBMCs are described, which range from 300 to 640 g for 7–10 min for the first wash step and from 300 to 470 g for 7–10 min for the second wash step [26,41,51,53–55].

Finally, the amount of PBMCs used varied slightly, as we used $2.5 \times 10^5$ PBMCs/well, and according to the LymeSpot assay, $2.0 \times 10^5$ PBMCs/well should have been used. A higher number of PBMCs per well could result in increased numbers of SFCs, as the use of more PBMCs results in more antigen-presenting cells and more T cells that could become activated after stimulation with the Borrelia antigens.

The results of a comparative pilot experiment that we performed in which we assessed the influence of the deviations discussed above supported that these deviations from the recommended protocol are not critical as such (Supporting information, Data S4). Hence, the conclusion stands that both ELISpot assays cannot help to diagnose active LNB.

Probably some of the most critical steps that influence the performance of an ELISpot assay are the time between venipuncture and PBMC isolation, the time between PBMC isolation and incubation of the assay and the (overnight) incubation time of the assay [56,57]. In this study, these times were all within the limits as described in the LymeSpot protocol, with the exception of the time between venipuncture and PBMC isolation, which was prolonged for various cases. A prolonged time between venipuncture and PBMC isolation is known to decrease the PBMC viability [56]. To compensate for this, for those cases for which the time between venipuncture and PBMC isolation was prolonged (8–32 h), we performed a T-Cell Xtend step prior to PBMC isolation. This T-Cell Xtend step has proved not to be detrimental to the PBMC yield and the ELISpot performance [26,27,58].

No data are provided in the instruction manual of the LymeSpot assay with regard to the diagnostic performance of this assay. To our knowledge, this is the first study that has investigated the diagnostic capacity of the LymeSpot assay for the diagnosis of active LNB. The diagnostic performance of the LymeSpot assay for other manifestations of LB has not yet been investigated thoroughly and remains unclear. Hopefully, more validation studies will be performed which will include other manifestations of LB, as well as follow-up studies to understand more clearly the diagnostic potential for treatment monitoring.

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Disclosures

None.

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Supporting Information

Additional supporting information may be found in the online version of this article at the publisher’s web site:

Table S1. Comparison of the protocols used for the evaluation of the in-house Borrelia ELISpot assay and the LymeSpot assay.

Fig. S2. Interpretation scheme of the LymeSpot assay according to the manufacturer.

Table S3. Overview of the study populations used in the three studies.

Data S4. A pilot study to investigate the influence of three different protocols for the isolation of peripheral blood mononuclear cells (PBMCs) on the LymeSpot results.