An ultra-high-density protein microarray for high throughput single-tier serological detection of Lyme disease

Vasanth Jayaraman1,4, Karthik Krishna1, Yuanyuan Yang2, Karenah J. Rajasekaran2, Yuzheng Ou2, Tianhao Wang3, Kang Bei1, Hari Krishnan Krishnamurthy1,4,*, John J. Rajasekaran1, Alex J. Rai3 & Daniel A. Green3*

Current serological immunoassays have inherent limitations for certain infectious diseases such as Lyme disease, a bacterial infection caused by *Borrelia burgdorferi* in North America. Here we report a novel method of manufacturing high-density multiplexed protein microarrays with the capacity to detect low levels of antibodies accurately from small blood volumes in a fully automated system. A panel of multiple serological markers for Lyme disease are measured using a protein microarray system, Lyme Immunochip, in a single step but interpreted adhering to the standard two-tiered testing algorithm (enzyme immunoassay followed by Western blot). Furthermore, an enhanced IgM assay was supplemented to improve the test’s detection sensitivity for early Lyme disease. With a training cohort (n = 40) and a blinded validation cohort (n = 90) acquired from CDC, the Lyme Immunochip identified a higher proportion of Lyme disease patients than the two-tiered testing (82.4% vs 70.6% in the training set, 66.7% vs 60.0% in the validation set, respectively). Additionally, the Immunochip improved sensitivity to 100% while having a lower specificity of 95.2% using a set of investigational antigens which are being further evaluated with a large cohort of blinded samples from the CDC and Columbia University. This universal microarray platform provides an unprecedented opportunity to resolve a broad range of issues with diagnostic tests, including multiplexing, workflow simplicity, and reduced turnaround time and cost.

Serological tests, which can be used to detect and characterize the antibody response to infection, are primarily formatted as immunoassays, including enzyme immunoassays (EIA), enzyme-linked immunosorbent assays (ELISA), Western blot (WB), immunoblot (IB), etc1-2. However, existing immunoassays have three fundamental limitations that have impeded their applications in disease diagnosis. First, conventional immunoassays are limited by their level of multiplexing, defined as the maximum number of antibodies that can be detected in a single run3. Second, most immunoassays have limited sensitivity for detection of antibodies at low levels, which can impede their utility in early stage diagnosis or in individuals with compromised immune systems with low antibody production4. Third, there is a lack of robust and flexible technology which can readily accept the addition of novel antigen biomarkers that have been validated for clinical use5.

Lyme disease, a bacterial infection caused primarily by *Borrelia burgdorferi* and a few cases by *Borrelia mayonii* in North America, is increasing in prevalence and public health importance, but its diagnosis can be impeded by the lack of accurate laboratory tests, particularly for early detection5-6. While early localized disease is usually accompanied by a characteristic erythema migrans (EM) rash, up to 20–30% of patients may lack this finding7-8 and if left untreated, bacteria can disseminate to cause nervous system involvement and/or cardiac conduction abnormalities9-10. Late disease can present as arthritis and/or a wide spectrum of nervous system manifestations11. In the United States, approximately 300,000 people are estimated to contract Lyme disease each year, but only 42,743 confirmed and probable cases of Lyme disease were reported to Centers for Disease Control and Prevention (CDC) in 2017, an increase of 17% from 201612.

1Vibrant Sciences LLC., San Carlos, CA, USA. 2Vibrant America LLC., San Carlos, CA, USA. 3Department of Pathology and Cell Biology, Columbia University Irving Medical Center, New York, NY, USA. 4These authors jointly supervised this work: Vasanth Jayaraman and Hari Krishnan Krishnamurthy. 5email: hari@vibrantsci.com; dag2149@cumc.columbia.edu
Assessment of the antibody response to *B. burgdorferi* infection has been the mainstay of laboratory confirmation of Lyme disease for over 20 years. A two-tiered serology algorithm, which utilizes a first-tier EIA followed by a second-tier WB or IB assay, was recommended in the United States and Europe (Fig. 2A)\(^\text{13,14}\). Contemporary EIAs have been developed using select recombinant proteins and/or select synthetic peptides from immunodominant regions within proteins that are specific to and conserved among the *B. burgdorferi* sensu lato (sl) complex members. WB analysis has been required to supplement positive EIA tests owing to its higher specificity and the ability to detect multiple protein antigens. However, blot-based testing is known to cause false positive results related to over-reading of weak bands in its gel matrix that affect quality of testing in Lyme disease, especially for IgM interpretation\(^\text{15}\). The two-tiered testing is inherently labor-intensive and uses subjective criteria for interpretation of results that may impact reproducibility with increased turnaround time and added complexity and cost\(^\text{16}\).

Previously, we developed a silicon-based peptide microarray platform utilizing a high-volume manufacturing and computational process derived from semiconductor industrial design. This novel technology has been validated in identification of highly predictive peptide antibody biomarkers in diagnosis and monitoring of Celiac disease\(^\text{17,18}\). In this study, we introduce a multiplex microarray of *Borrelia burgdorferi* recombinant antigens, wherein the proteins are physically separated from each other, to detect low levels of serological markers for Lyme disease from small blood volumes in a fully automated workflow (Fig. 2B). We explore here a novel testing methodology using a training cohort and blinded clinical samples from the CDC. With a functionalized high-density substrate, enhanced IgM assay, and a set of investigational markers, we aimed to improve both the analytical and clinical performance characteristics in Lyme disease testing.

### Materials and methods

#### Patient samples. Two cohorts of serum samples acquired from CDC Lyme serum repository (LSR) were employed\(^\text{19}\). The training cohort contained 40 sera and the blinded validation cohort contained 90 sera. These serum samples were from Lyme disease positive patients at various stages with different sequelae, and negative controls including healthy controls and patients with other diseases. The composition of both cohorts is presented in Table 3. The informed consent and Institutional Review Board approval for testing these samples was granted during the establishment of LSR by CDC. Testing of reference sera was performed by laboratory personnel at Vibrant America Clinical Labs (CLIA and CAP accredited facility) without prior knowledge of the expected results in the same manner as clinical samples from Lyme disease patients. The CDC provided EIA results, detectable bands from WBs, WB interpretation by CDC criteria, and two-tiered testing results. CDC utilized VIDAS Lyme IgM and IgG polyvalent assay by bioMérieux, Inc for the EIA testing and MarDx Diagnostics IgM and IgG immunoblotting assays for the WB testing. The 142 healthy negative sera samples used to establish cut-off (CU) values was collected and tested under IRB exemption (work order #1-1098539-1) determined by the Western Institutional Review Board (WIRB) to use de-linked and de-identified human specimen and medical data for research purposes. The upper 97.5th percentile for each analyte was set to 10 CU and the upper 99th percentile was set to 20 CU (Supplemental Dataset). All methods used in this study were carried out in accordance with relevant guidelines and regulations.

#### Wafer substrate preparation. Prime grade 300 mm silicon wafers with p-type boron, (1 0 0) orientation, resistivity of 1 to 5 Ω cm\(^{-1}\), and 725 μm thickness were deposited with 100 nm thermal oxide by dry oxidation at 1,000 °C in a furnace under pure oxygen atmosphere for two hours. Wafers were copiously washed with deionized water for five minutes and spin coated with a solution containing 1% (vol/vol) of 3-aminopropyltriethoxysilane (APTES) in N-methylpyrrolidone (NMP) and left at room temperature for 15 min. Curing of the wafers was done at 120 °C for 60 min under N\(_2\) atmosphere and humidity-controlled environment. A copolymer solution containing poly(L-lysine) and poly(lactic acid) was coated on the wafer and reacted for 24 h to generate a high binding surface capable of immobilizing proteins via passive adsorption through hydrophobic interactions with lysine polymer and covalent coupling with the lactic acid polymer (structures shown in Fig. 1B). The functionalized wafers were normally used within one week of preparation.

#### Protein attachment. The following antigens were included in the panel: *Borrelia burgdorferi* VlsE1, *Borrelia burgdorferi* p18, *Borrelia burgdorferi* p23, *Borrelia burgdorferi* p28, *Borrelia burgdorferi* p30, *Borrelia burgdorferi* p31, *Borrelia burgdorferi* p34, *Borrelia burgdorferi* p39, *Borrelia burgdorferi* p41, *Borrelia burgdorferi* p45, *Borrelia burgdorferi* p58, *Borrelia burgdorferi* p66, *Borrelia burgdorferi* p93. Figure 1C shows *Borrelia burgdorferi* p23 as an example for the protein immobilization process. The recombinant antigens were expressed in E. coli bacterial cells using full length cDNA coding for the respective antigens fused with a hexa histidine purification tag. The capture antigens were incubated on the wafer at a concentration of 1.0 μg/ml and reacted for 24 h at 4 °C. Excess unbound antigens were removed by extensive washing with aqueous phosphate buffer and the unreacted substrate was quenched with a blocking solution containing BSA and glycine. Each wafer was patterned with a layout of 44 microchips.

#### Pillar plate assembly. The pillar plate assembly process is illustrated in Fig. 1d–g. The individual wafers for each antigen were then diced into 0.7 × 0.7 mm\(^2\) microchips using a stealth dicing process, as previously described\(^\text{21}\). The diced wafers were picked and placed onto individual carrier tapes using a standard die sorting system. The carrier tapes were loaded onto a high-throughput surface mount technology (SMT) component placement system. The microchips were then picked and placed onto 24-pillar plates with each pillar containing a layout of 44 microchips.
Immunochip assay. Serum samples were probed using 1:20 dilution on the pillar plate and reacted for one hour at room temperature as previously described. The plate was then washed with Tris-buffered saline with Tween 20 (TBST) (Amresco) buffer 3 × 5 min each. The plate was incubated with the secondary antibody (1:2000 dilution of Goat Anti-Human IgG HRP and Goat Anti-Human IgM HRP individually) for one hour at room temperature. The plates were washed with TBST buffer followed by washing with DI Water. The plates were finally dried completely before adding chemiluminescent substrate (Lumi-Phos HRP from Lumigen), per manufacturer recommendations and scanned for five minutes on a standard Chemiluminescence Imager. For the Enhanced IgM Assay, the serum was pre reacted with Goat anti-human IgG Fc fragment prior to the remaining assay steps to increase the sensitivity of IgM detection.

Data analysis. The generated images are analyzed using the in-house reporter software as described previously. Briefly, the chemiluminescent signals are converted into intensity plots which was compared to the thresholds established for each antigen to determine sero-positivity.

Results

Lyme immunochip platform. The main components of the Lyme Immunochip platform include multiple silicon-based 0.70 × 0.70 mm² microchips that are laser diced from antigen-immobilized wafers, a customized 24-well compatible plate containing 24 pillars, each containing 44 microchips that are picked and placed into a multiplex microarray assembly, and a high-resolution imager capable of simultaneously detecting chemiluminescent signals from labeled antigen–antibody reactions at each microchip throughout the multiplex microarray (Fig. 1). Each chip can be considered analogous to an individual band in a Western blot; however, the proteins are physically separated eliminating cross-reactive issues seen in blot-based assays for proteins with similar mass (Fig. 2C).

Multiplex antibody detection. Detection of multiplex antibodies is based on a chemiluminescent immunoassay. In contrast to two-tiered testing by EIA and WB, the multiplex test on Lyme Immunochip requires only 25.0 μL of serum specimen. The entire process including sample dilution, multi-step incubation, and multi-solution washing is programmed into liquid handlers. Coupled with automatic detection of analytes and data acquisition, the Lyme Immunochip has the capacity to simultaneously assay 192 individual specimens within 2 h. An in-house reporter software is designed to extract the raw chemiluminescent signals for each probe and convert them into intensity plots after quantile normalization, background, and spatial correction. Compared with traditional two-tiered testing (Fig. 2A), automatic detection of multiple antigens in this high-throughput manner (Fig. 2B) has the potential to dramatically shorten turnaround time, reduce the cost of labor and instru-
Figure 2. (A) Standard two-tiered testing algorithm recommended by CDC. (B) Vibrant Lyme Immunochip testing algorithm. (C) Different bands on a typical Western blot / Immunoblot are tested one protein per microchip using multiple microchips per pillar.

Table 1. Composition of the Vibrant’s Lyme Immunochip and interpretation with the standard two-tiered testing algorithm.

| Species          | Antigen                  | Antibody |
|------------------|--------------------------|----------|
| 1st Tier         |                          |          |
| B. burgdorferi   | VlsE1                     | IgM/IgG  |
| 2nd Tier         |                          |          |
| B. burgdorferi   | p23(OspC), p39(BmpA), p41(Fla) | IgM     |
| B. burgdorferi   | p18(DBpB), p23(OspC), p28(Oms28), p30, p39(BmpA), p41(Fla), p45, p58(OppA-2), p66(Oms66), p93 | IgG     |
| For research purpose |                       |          |
| B. burgdorferi   | p18(DBpB), p28(Oms28), p30, p31(OspA), p34(OspB), p45, p58(OppA-2), p66(Oms66), p93 | IgM     |
| B. burgdorferi   | p31(OspA), p34(OspB)      | IgG     |
To further improve the accuracy for diagnosis of early Lyme disease, an in-house enhanced IgM assay was developed and utilized. IgM antibodies are the first type of antibody produced by the immune system upon infection. An IgM antibody test is often used in the early stages of Lyme disease diagnosis, typically within the first month of infection. The test evaluates for the presence of IgM antibodies against B. burgdorferi, the bacteria that causes Lyme disease. By detecting IgM antibodies early in the infection, the test can help identify individuals in the acute phase of Lyme disease, allowing for prompt treatment and management.

Enhanced IgM assay.

Table 2. Analytical performance of the Lyme Immunochip.

| Antibody       | Coefficient of variation for repeatability (%) | Coefficient of variation for reproducibility (%) | Analytical sensitivity |
|----------------|-----------------------------------------------|-----------------------------------------------|-----------------------|
|                | Sample 1 | Sample 2 | Sample 3 | Sample 1 | Sample 2 | Sample 3 | LoB (CU) | LoD (CU) |
| B. Burgdorferi VlsE1 IgM | 2.87     | 3.33     | 5.30     | 6.95     | 3.77     | 1.77     | 0.3       | 0.8      |
| B. Burgdorferi VlsE1 IgG | 4.14     | 1.92     | 1.93     | 5.28     | 4.70     | 1.26     | 0.3       | 0.7      |
| B. Burgdorferi p23 IgM | 4.79     | 2.11     | 3.84     | 1.94     | 1.57     | 2.94     | 0.3       | 0.8      |
| B. Burgdorferi p39 IgM | 3.39     | 3.72     | 4.11     | 3.64     | 1.73     | 2.44     | 0.4       | 0.8      |
| B. Burgdorferi p41 IgM | 3.93     | 2.93     | 2.65     | 1.73     | 6.34     | 1.93     | 0.1       | 0.7      |
| B. Burgdorferi p18 IgG | 4.04     | 1.85     | 2.63     | 2.09     | 2.05     | 2.04     | 0.4       | 0.8      |
| B. Burgdorferi p23 IgG | 3.10     | 9.52     | 2.21     | 4.72     | 2.03     | 1.72     | 0.3       | 0.6      |
| B. Burgdorferi p28 IgG | 1.56     | 5.08     | 1.56     | 2.15     | 3.92     | 7.14     | 0.5       | 0.8      |
| B. Burgdorferi p30 IgG | 1.53     | 4.42     | 3.59     | 2.73     | 3.92     | 8.13     | 0.4       | 0.6      |
| B. Burgdorferi p39 IgG | 1.26     | 8.59     | 2.59     | 2.37     | 2.32     | 2.89     | 0.3       | 0.8      |
| B. Burgdorferi p41 IgG | 1.91     | 1.68     | 1.29     | 2.33     | 5.67     | 2.28     | 0.2       | 0.5      |
| B. Burgdorferi p45 IgG | 1.49     | 1.62     | 2.29     | 4.69     | 4.40     | 3.65     | 0.4       | 0.5      |
| B. Burgdorferi p58 IgG | 1.57     | 2.34     | 4.58     | 5.30     | 4.43     | 1.47     | 0.2       | 0.7      |
| B. Burgdorferi p66 IgG | 4.15     | 2.18     | 8.69     | 4.80     | 2.67     | 8.37     | 0.1       | 0.6      |
| B. Burgdorferi p93 IgG | 1.82     | 4.00     | 1.75     | 6.44     | 1.61     | 4.61     | 0.2       | 0.7      |
the immune system in response to an infection. IgM antibodies only comprise 5% to 10% of all the antibodies present in circulation, whereas IgG antibodies, which are the most abundant immunoglobulin, comprise about 75% to 80%. In order to enhance sensitivity and specificity of the IgM assay, we reduced assay interference by removing a major portion of the abundant IgG antibodies and other non-specific proteins from the serum while detecting the IgM antibodies. A purified goat anti-human (GAH) IgG Fc fragment was incubated with serum for the removal of human IgG prior to testing for specific IgM antibodies. In the training set, the enhanced IgM assay further improved the sensitivity to 70.0% for early Lyme patients while the performance was equivalent for the rest of the disease/control samples as shown in Table 3. Samples captured by enhanced IgM but not standard IgM are shown in Supplementary Figure S1. There were also no additional false positives with the enhanced IgM assay as compared to the standard IgM assay as seen in Fig. 3.

Blinded clinical sample validation. To validate the clinical performance of the Lyme Immunochip, a cohort of blinded samples (N = 90) acquired from the CDC LSR was tested and interpreted by criteria achieved via the training set. After incorporating the enhanced IgM assay, the single-tier Immunochip was compared with standard two-tiered serologic testing. Whereas the Immunochip identified 9 out of 18 early Lyme disease patients (50.0%) at acute and convalescent phases, the standard two-tiered testing identified only 7 of them (38.9%). The clinical performance by both methods were identical towards later-stage Lyme disease, including neurologic Lyme disease (75.0%), cardiac Lyme disease (100.0%), and Lyme arthritis (100.0%). The specificity levels achieved by this single-tier assay was identical to that of the two-tiered testing (96.7%). The heat map in Fig. 3 shows antibody-binding intensity data differentiating samples seropositive for Lyme disease from healthy

| Table 3. Clinical performance of the Lyme Immunochip in the training and validation cohorts. |
|---|
| **Cohort and subject type** | **n** | **Vibrant Lyme Immunochip** | **Standard two-tiered testing** | **Enhanced IgM** | **Standard** |
| **Sensitivity (%)** | **Specificity (%)** | **Sensitivity (%)** | **Specificity (%)** | **Sensitivity (%)** | **Specificity (%)** |
| **Training cohort** | **40** | | | | |
| Early Lyme disease—acute/convalescent | 10 | 70.0 | 60.0 | 50.0 | |
| Neurologic Lyme | 3 | 100.0 | 100.0 | 100.0 | |
| Lyme arthritis | 4 | 100.0 | 100.0 | 100.0 | |
| Total positive: | 17 | 82.4 | 76.5 | 70.6 | |
| Fibromyalgia | 2 | 100.0 | 100.0 | 100.0 | |
| Rheumatoid arthritis | 2 | 100.0 | 100.0 | 100.0 | |
| Mononucleosis | 2 | 100.0 | 100.0 | 100.0 | |
| Syphilis | 2 | 100.0 | 100.0 | 100.0 | |
| Multiple sclerosis | 2 | 100.0 | 100.0 | 100.0 | |
| Severe periodontitis | 2 | 100.0 | 100.0 | 100.0 | |
| Healthy endemic | 7 | 100.0 | 100.0 | 100.0 | |
| Healthy non-endemic | 4 | 100.0 | 100.0 | 100.0 | |
| Total negative: | 23 | 100.0 | 100.0 | 100.0 | |
| **Validation cohort** | **90** | | | | |
| Early Lyme disease—acute/convalescent | 18 | 50.0 | 38.9 | 38.9 | |
| Neurologic Lyme | 4 | 75.0 | 75.0 | 75.0 | |
| Cardiac Lyme | 2 | 100.0 | 100.0 | 100.0 | |
| Lyme arthritis | 6 | 100.0 | 100.0 | 100.0 | |
| Total positive: | 30 | 66.7 | 60.0 | 60.0 | |
| Fibromyalgia | 6 | 100.0 | 100.0 | 100.0 | |
| Rheumatoid arthritis | 6 | 100.0 | 100.0 | 100.0 | |
| Mononucleosis | 6 | 100.0 | 100.0 | 100.0 | |
| Syphilis | 6 | 66.7 | 66.7 | 83.3 | |
| Multiple sclerosis | 6 | 100.0 | 100.0 | 100.0 | |
| Severe periodontitis | 6 | 100.0 | 100.0 | 100.0 | |
| Healthy endemic | 12 | 100.0 | 100.0 | 100.0 | |
| Healthy non-endemic | 12 | 100.0 | 100.0 | 100.0 | |
| Total Negative | 60 | 96.7 | 96.7 | 96.7 | |
| Complete panel with established markers | 130 | 72.3 | 97.6 | 66.0 | 97.6 | 63.8 | 97.6 |
| Complete panel with investigational markers | 130 | 100.0 | 95.2 | 74.5 | 94.0 | NA | NA |
controls and disease controls. The enhanced IgM assay was able to recognize two early Lyme disease samples at convalescent phase that were missed by the standard IgM assay. Cardiac Lyme disease, neurologic Lyme disease and Lyme arthritis exhibited higher-intensity binding by the IgG assay as compared to the IgM assays (Fig. 3). The Immunochip demonstrated sensitivity and specificity of 66.7% and 96.7% compared with those of CDC’s 60.0% and 96.7% in the blinded validation cohort. A set of investigational markers including p31 and p34 (Table 1) was also separately evaluated with interpretative criteria requiring binding of the first-tier antigen (VlsE1) plus any two second-tier antigens. While these investigational markers demonstrated higher sensitivity (100%) than the standard two-tiered antigen panel, they were also associated with lower specificity (95.2%) and require further evaluation with expanded sample sets. As shown in Fig. 3, the improvement in sensitivity was almost exclusively due to the inclusion of p31 and p34 in the enhanced IgM assay, which was particularly useful for detecting patients with early Lyme Disease.

Discussion
Here, we present the first ultra-high-density microarray platform to rapidly and accurately detect a panel of well-characterized Lyme disease serological markers. The Lyme Immunochip applies the standard two-tiered testing algorithm criteria, but all markers are detected in a single run, which substantially improves turnaround time, reduces the cost of labor and instrument, and eliminates the need of manual handling and subjective interpretation of the WB or IB test results.

Multiplexing of proteins has been previously performed using spotted arrays or WB/IB26. Although some of these platforms have been used in diagnostic testing and life science research, they have important limitations. Protein spraying/spotting techniques are inconsistent due to variation in the volume of reagents taken up and dispensed owing to differences in reagent viscosity27. “Missing spots” or “merged spots” result from uneven surfaces, which hinders consistent contact between the printers/spotters and the chip surface28. Blot-based techniques additionally require skilled technologists to interpret bands, which leads to variability in interpretation, especially for weak bands29. Although automated readers have been reported for immunoblots, manufacturing technology of existing multiplex platforms are not capable of high throughput testing and mass production, which impacts their clinical utility at scale30,31.

To address these issues, we fundamentally innovated on the technology at every stage, including chip manufacturing, surface functionalization, array assembly, analyte detection, and data acquisition in the Vibrant Protein Microarray platform. The Vibrant pillar plate can incorporate 100 probe-immobilized microchips (in its current configuration) at each pillar, i.e., 100 types of biomarkers can potentially be detected for 24 samples in one pillar plate within a 1.0 cm² pillar. Further compaction of the microchip can increase the number of biomarkers on each pillar plate and enhance the total throughput. For example, reducing the microchip size from 0.70 × 0.70 mm² to 0.35 × 0.35 mm², which is feasible (data not shown), would quadruple the capacity to 400 probes per pillar. Importantly, given that Ixodes ticks transmit other human pathogens such as Anaplasma, Babesia, and Ehrlichia, this ultra-high level of multiplexing allows for the further development of panels that can simultaneously assess for multiple tickborne diseases from a single, low-volume blood sample, with the added flexibility to incorporate emerging pathogens that become more prevalent over time.

Clinically the Lyme Immunochip showed higher overall sensitivity (72.3%) than standard two-tier testing (63.8%) (Table 3). VlsE1 and OspC were the two most sensitive markers observed on our Lyme Immunochip (a complete list of individual B. burgdorferi antigens’ clinical performance is shown in Figure S1). VlsE1 is composed...
of alternating variable (VR1 to VR6) and invariable (IR1 to IR6) regions, among which IR6 has been identified as an immunodominant epitope. OspC is the second most-commonly targeted antigen in diagnostic testing and has an important role in transmission of Lyme disease-associated Borrelia from the tick vector to the mammalian host. OspC is available for immune stimulation sooner than VlsE because it is expressed on the surface of Borrelia burgdorferi before infection.

These sensitivity gains did not come at the cost of specificity, which was equivalently high for both the Lyme Immunochip and standard two-tier testing (97.6% each). Test specificity is a paramount concern for Lyme disease assays to mitigate the risk of false positives in low prevalence settings, and we therefore designed the Lyme Immunochip to utilize the same highly specific markers and stringent interpretation criteria used in the standard two-tier algorithm. The most specific markers in the panel were p58 and p39. The function of p58 is suggested to be involved in the transmembrane transport of solutes and but it shares a functional domain usually found in periplasmic oligopeptide-binding proteins. B. burgdorferi basic membrane protein A (BmpA), also known as p39, localizes to the bacterium’s outer membrane and binds to laminin in the host's extracellular matrix. The high specificity observed for these two markers is in accordance with previous reports but their exact functions are still elusive. Recently, CDC recommended a modified two-EIA algorithm as an alternative to Western blot confirmation. While this approach allows for the accurate diagnosis of most straightforward cases with lower cost and faster turnaround time than Western blot, knowledge of which bands/antigens are reactive can aid in the interpretation of challenging cases, help differentiate new vs. past infections, and also measure the expansion of immune response over time. The Lyme Immunochip can provide the same detailed information as Western blot, but with a faster turnaround time that is offered by a two-EIA approach. The test report can therefore include detailed antigen reactivity along with an overall positive/negative interpretation according to CDC guidelines and duration of symptoms (Fig. 2).

As the known repertoire of B. burgdorferi surface and secreted proteins expands over time, additional markers can be easily incorporated into the platform, enabling further performance improvements. In a previous study, a ten-antigen panel was selected from 62 B. burgdorferi surface proteins and synthetic peptides, which yielded 87.5% sensitivity for early Lyme disease by scoring the presence of any two markers as a positive result. Based on a similar criterion, our preliminary study using our current protein microarray with all investigational antigens (as listed in Table 1) raised the sensitivity to 100.0% while lowering specificity to 95.2% for the two cohorts altogether; further validation with a large blinded cohort is underway. In addition, we are actively investigating previously unappreciated epitopes of highly immunogenic antigens that are targeted by the immune system using a peptide microarray. Generating an ultra-high-density peptide microarray with these additional epitopes may further improve clinical performance, identify disease subsets, and also provide fundamental insights into the pathogenesis of Lyme disease, including post-treatment Lyme disease syndrome.

In conclusion, we demonstrate that the Lyme Immunochip is a valid alternative to standard two-tiered testing, with the capability to characterize the antibody profile of Lyme disease patients with enhanced accuracy and reduced turnaround time. The advances in semiconductor methods and the generation of high-throughput microarrays for creating antigen diversity provides the potential to improve the diagnostic accuracy of immunoassays. The novel protein microarray method described herein provides three major advantages over existing testing methodologies: (1) an ultra-high-density array surface, (2) higher reproducibility, and (3) improved throughput. This platform permits simultaneous detection of a large number of candidate biomarkers from low volume samples with reduced cost and turnaround time, allowing for both multiplex diagnostic testing as well as biomarker discovery and validation. This ultra-high-density microarray has the potential to be an accurate, rapid, economic, and universal diagnostic testing platform.
16. Wormser, G. P. et al. Comparative cost-effectiveness of two-tiered strategies for serodiagnosis of Lyme disease with noncutaneous manifestations. J. Clin. Microbiol. 51, 4045–4049 (2013).
17. Choung, R. S. et al. Determination of B-cell epitopes in patients with celiac disease: Peptide microarrays. PLoS ONE 11, e0147777 (2016).
18. Choung, R. S. et al. Synthetic neoepitopes of the transglutaminase-deamidated gliadin complex as biomarkers for diagnosing and monitoring celiac disease. Gastroenterology 156, 582–591 (2019).
19. Centers for Disease Control and Prevention. Updated CDC Recommendation for Serologic Diagnosis of Lyme Disease. www.cdc.gov/mmwr/volumes/68/ww/mm6832a4.htm#contribAff. Accessed September 23, 2019.
20. Kumaran, D. et al. Crystal structure of outer surface protein C (OspC) from the Lyme disease spirochete, Borrelia burgdorferi. EMBO J. 20, 971–978 (2001).
21. Krishnamurthy et al. Antibody profiling and prevalence in the US population during the SARS-CoV2 Pandemic. medRxiv 2020.04.29.20085068.
22. Bruckbauer, H. R., Preac-Mursic, V., Fuchs, R. & Wilske, B. Cross-reactive proteins of Borrelia burgdorferi sensu lato) and Borrelia burgdorferi as a marker for infection in experimentally and naturally inoculated animals. J. Clin. Microbiol. 38, 208–215 (2000).
23. Ang, C. W. et al. Large differences between test strategies for the detection of anti-Borrelia antibodies are revealed by comparing eight ELISAs and five immunoblots. Eur. J. Clin. Microbiol. Infect. Dis. 30, 1027–1032 (2011).
24. Schriever, M. E. et al. Collection and characterization of samples for establishment of a serum repository for Lyme disease diagnostic test development and evaluation. J Clin Microbiol 52, 3755–3762 (2014).
25. Terry, W. D. & Fahey, J. L. Subclasses of human gamma-2-globulin based on differences in the heavy polypeptide chains. Science 146, 400–401 (1964).
26. Gupta, S., Manubhai, K. P., Kulkarni, V. & Srivastava, S. An overview of innovations and industrial solutions in Protein Microarray Technology. Proteomics 16, 1297–1308 (2016).
27. Kambhampati, D. Protein Microarray Technology (Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, 2003).
28. Voskuhl, J., Brinkmann, J. & Jonkheijm, P. Advances in contact printing technologies of carbohydrate, peptide and protein arrays. Curr. Opin. Chem. Biol. 18, 1–7 (2014).
29. Dressler, F., Whalen, J. A., Reinhardt, B. N. & Steere, A. C. Western blotting in the serodiagnosis of Lyme disease. J. Infect. Dis. 167, 392–400 (1993).
30. Phizicky, E., Bastaens, P. I., Zha, H., Snyder, M. & Fields, S. Protein analysis on a proteomic scale. Nature 422, 208–215 (2003).
31. MacBeath, G. & Schreiber, S. L. Printing proteins as microarrays for high-throughput function determination. Science 289, 1760–1763 (2000).
32. Chandra, A., Latov, N., Wormser, G. P., Marques, A. R. & Alaedini, A. Epitope mapping of antibodies to VlsE protein of Borrelia burgdorferi in post-Lyme disease syndrome. Clin. Immunol. 141, 103–110 (2011).
33. Padula, S. J., Dias, F., Sampieri, A., Craven, R. B. & Ryan, R. W. Use of recombinant OspC from Borrelia burgdorferi for serodiagnosis of early Lyme disease. J. Clin. Microbiol. 32, 1733–1738 (1994).
34. Hauser, U., Lehnert, G. & Wilske, B. Diagnostic value of proteins of three Borrelia species (Borrelia burgdorferi sensu lato) and implications for development and use of recombinant antigens for serodiagnosis of Lyme borreliosis in Europe. Clin. Diagn. Lab. Immunol. 5, 456–462 (1998).
35. Simpson, W. J., Burgdorfer, W., Schrumpf, M. E., Karstens, R. H. & Schwan, T. G. Antibody to a 39-kilodalton Borrelia burgdorferi antigen (P39) as a marker for infection in experimentally and naturally inoculated animals. J. Clin. Microbiol. 29, 236–243 (1991).
36. Steere, A. C., McHugh, G., Damle, N. & Sikand, V. K. Prospective study of serologic tests for Lyme disease. Clin. Infect. Dis. 47, 188–195 (2008).
37. Fawcett, P. T. et al. Detection of antibodies to the recombinant P39 protein of Borrelia burgdorferi using enzyme immunoassay and immunoblotting. J. Rheumatol. 20, 734–738 (1993).
38. https://www.captoadayonline.com/no-blot-testing-charts-new-course-for-lyme-dx/. Accessed February 12, 2020.
39. Lahey, L. J. et al. Development of a multitarget panel for improved detection of Borrelia burgdorferi Infection in early Lyme disease. J. Clin. Microbiol. 53, 3834–3841 (2015).

Acknowledgements

The authors thank Christopher Sexton and the staff of CDC for their distribution of the Lyme serum samples. Research reported in this publication was supported by the Vibrant America LLC. The content is solely the responsibility of the authors and does not necessarily represent the official views of the CDC. We also would like to thank Miaomiao Sun for generating the Figs. 1 and 2 illustrations.

Author contributions

H.K.K., J.J.R., D.A.G. designed the study, K.K., Y.Y. collected the clinical data, V.J., H.K.K., Y.Y. wrote the manuscript; K.K., Y.O., T.W. collected and tested the samples; Y.Y., K.K., K.E.R. analyzed the data; Y.O., K.K. carried out the experiments, D.A.G., J.J.R., H.K.K. revised the manuscript critically for important intellectual content, KB developed the Vibrant Imaging Software. All the authors read and approved the final manuscript.

Competing interests

All of the authors listed in this paper are employees of Vibrant Sciences or Vibrant America, with the exception of AJR and DAG, who are academic collaborators and consultants to Vibrant Sciences.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-75036-2.
Correspondence and requests for materials should be addressed to H.K.K. or D.A.G.
Reprints and permissions information is available at www.nature.com/reprints.
Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
