Monoclonal Antibodies for the Diagnosis of *Borrelia crocidurae*

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Abstract. Relapsing fever borreliae, produced by ectoparasite-borne *Borrelia* species, cause mild to deadly bacteremia and miscarriage. In the perspective of developing inexpensive assays for the rapid detection of relapsing fever borreliae, we produced 12 monoclonal antibodies (MAbs) against *Borrelia crocidurae* and characterized the two exhibiting the highest titers. P3A10 MAb reacts with the 35.6-kDa flagellin B (flaB) of *B. crocidurae* while P6D9 MAb recognizes a 35.1-kDa variable-like protein (Vlp) in *B. crocidurae* and a 35.2-kDa Vlp in *Borrelia duttonii*. Indirect immunofluorescence assay incorporating relapsing fever and Lyme group borreliae and 11 blood-borne organisms responsible for fever in West Africa confirmed the reactivity of these two MAbs. Combining these two MAbs in indirect immunofluorescence assays detected relapsing fever borreliae including *B. crocidurae* in ticks and the blood of febrile Senegalese patients. Both antibodies could be incorporated into inexpensive and stable formats suited for the rapid point-of-care diagnosis of relapsing fever. These first-ever MAbs directed against African relapsing fever borreliae are available for the scientific community to promote research in this neglected field.

INTRODUCTION

Relapsing fever borreliae are arthropod-borne pathogens causing mild to deadly spirochetemia, most commonly resulting in malaria-like symptoms.¹ In Africa, cultured representatives include tick-borne *Borrelia crocidurae*, *Borrelia duttonii*, and *Borrelia hispanica*, transmitted by *Ornithodoros* soft ticks, and louse-borne *Borrelia recurrentis*.² Recently, we detected, using molecular tools, a new *Borrelia* sp. named *Candidatus Borrelia algerica* in the blood of febrile patients in Algeria.³ Resource-consuming molecular methods are required to detect these pathogens in vectors and clinical specimens, which may not be routinely available in endemic regions,⁴ in addition to nonspecific direct microscopic examination. Microscopic examination is not able to distinguish between the various *Borrelia* species and molecular methods are not widely available, mainly used in a few reference centers not necessarily located in the endemic regions. However, prognosis of relapsing fever depends on the causative species, with the mortality rate being significantly higher for *B. recurrentis* than that for *B. crocidurae*.⁵ Indeed, part of the deadly mortality is due to Jarisch–Herxheimer reactions when treating patients infected by *B. recurrentis*; such a reaction has never been described with *B. crocidurae*.⁶ Development of inexpensive assays for the rapid detection of relapsing fever borreliae is therefore warranted. In this perspective, a few antibodies have been made using new world *Borrelia hermsii* and *Borrelia afzelii* (for the Lyme disease group) antigens, which may eventually cross-react with some African borreliae.⁷,⁸ and five murine monoclonal antibodies (MAbs): MAbs H5332 and H5TS specific for *Borrelia burgdorferi* outer surface protein A (OspA),⁹,¹⁰ MAbs H6831 and H614 specific for OspB,¹⁰ and MAb H9724 that reacts with a protein of the periplasmic flagella of the genus *Borrelia*.¹¹ However, no antibody has been developed using African relapsing fever borreliae as antigens. *Borrelia crocidurae* is the most common parasite of such relapsing fever borreliae in West Africa. We therefore produced and characterized MAbs against *B. crocidurae* with a view to incorporating them in a point-of-care laboratory test for rapid diagnosis.³

MATERIALS AND METHODS

Ethics statement. The *Ornithodoros sonrai* ticks studied here are not registered as endangered species. The study protocol was approved by the Steering Committee of the Institut Recherche et Développement (IRD) Special Program Évolution Climatique et Santé (Montpellier, France), reference project ATI-ECS-07-H/2002. As for the mammals, the study protocol was approved by Comité d’éthique de Marseille C2EA-14 and experimentation was performed according to the recommendations of reference project no. 60 12-11-2012. Regarding the human specimens, the study protocol was approved by the National Ethics Committee of Senegal Br 00081 04-06-12 and experimentation was performed according to the recommendations of reference projects SEN21/09 and SEN37/09.

*Borrelia* culture. *The* *B. crocidurae* Achema strain, (tick strain), *B. crocidurae* 03-02 strain, (clonial strain), *B. duttonii* Ly strain, *B. recurrentis* A1 strain, and *B. burgdorferi* B31 strain were grown at 32°C in a Barbour-Stoenner-Kelly-H medium (Sigma, Saint-Quentin-Fallavier, France) supplemented with 10% heat-inactivated rabbit serum (Eurobio, Courtaboeuf, France). Dark-field microscopic observation was performed to ensure the absence of any contaminant organisms and confirm the growth of the borreliae. To prepare antigens, broth inoculated with *B. crocidurae* was centrifuged at 14,000 × g for 10 minutes at 4°C; the pellet was washed twice with 5% phosphate-buffered saline (PBS) and Tween-20 (Euromedex, Souffelwayersheim, France), suspended in PBS and inactivated by incubation at 70°C for 1 hour.

Production of MAbs. To produce MAbs, 6-week-old female BALB/c mice were immunized by intraperitoneal inoculation of 10 μg of purified *B. crocidurae* Achema strain mixed with 100 μL of Imject Alum adjuvant (ThermoScientific, Courtaboeuf, France) (aluminum hydroxide and magnesium hydroxide mixture to stimulate the immune response for...
antibody production when incubated with immunogens v/v for 30 minutes). The mice were boosted three times at 2-week intervals, and then 5 days after the last injection, the spleens were sampled. Spleen cells were fused with myeloma cells (X63 Ag 8.653), then cultured in a supplemented Roswell Park Memorial Institute medium (Invitrogen, Cergy Pontoise, France) as previously described. Hybridoma supernatants were screened by an immunofluorometric assay. The purified B. crocidurae Achema isolate was spotted on glass slides and then fixed with methanol for 10 minutes. The hybridoma supernatants (200 µL) were deposited on spot and then incubated for 30 minutes at 37°C. The slides were washed twice with PBS containing 0.1% Tween for 5 minutes and with distilled water for 5 minutes. After drying, antibodies were tagged using a goat anti-mouse IgG conjugated to fluorescein isothiocyanate (FITC) (Immunotech, Marseille, France) at a 1:400 dilution in PBS. After washing, slides were dried, mounted with Fluoroprep (bioMérieux, Marcy l’Étoile, France), then observed under ultraviolet light using an epifluorescent Leica DM 2500 microscope (Leica, Saint-Jorioz, France) at a magnification of ×400. Sera from immunized mice were used as positive controls and those from healthy unexposed mice were used as negative controls. Cells of the wells secreted antibodies (determined by immunofluorescence assay with B. crocidurae) were cloned and subcloned two times. The iso-types of the MAbs were determined with an IsoStrip Mouse Monoclonal Isotyping Kit (Roche Diagnostic, Meylan, France) containing sensitized strips against IgG1, IgG2a, IgG2b, IgG3, IgA, and IgM. Then specificities of the MAbs were assessed by western immunoblotting.

Reactivity of MAbs. The MAbs produced were tested against B. duttonii, B. burgdorferi, B. recurrentis, and 11 blood-borne organisms responsible for fever in West Africa (Salmonella paratyphi, Streptococcus agalactiae, Streptococcus pneumoniae, Rickettsia felis, Plasmodium falciparum, Escherichia coli, CoxIELla burnetii, Bartonella quintana, Pseudomonas aeruginosa, Haemophilus influenzae, and Acinetobacter baumannii)3,13 by indirect immunofluorescence as described in the above section Production of MAbs. The sources and strains of the Borrelia species and other organisms used to screen hybridomas and test the reactivity of MAbs are presented in Table 1. To confirm the reactivity of these MAbs against B. crocidurae, we also tested a clinical strain B. crocidurae 03-02 isolated from a patient with relapsing fever in Senegal, and its genome sequenced in our laboratory.14

Gel electrophoresis and western blotting. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and western-blot analysis of Borrelia sp. were performed as previously described.15 Antigens were treated with proteinase K, and heat denaturation was performed at 100°C for 10 minutes. Borrelia crocidurae, B. burgdorferi, and B. duttonii whole-cell proteins resolved by 10% SDS-PAGE were visualized by silver staining. For one-dimensional western blotting, the gels were transferred to nitrocellulose and incubated with supernatants of MAbs P6D9 and P3A10, diluted at 1:100 and 1:500, respectively.

To prepare crude extracts for two-dimensional (2D) gel electrophoresis, purified bacteria were lysed by sonication in a rehydration solution (7 M urea, 2 M thiourea, 4% w/v 3-[(3-chloramidopropyl) dimethylammonio]-1-propanesulfonate hydrate) and centrifuged (10,000 × g, 20 minutes, 4°C) to remove cell debris and unbroken cells. The whole-cell protein extract was precipitated using the PlusOne 2-D Clean-Up Kit (GE Healthcare, Challfnt St. Giles, United Kingdom). The final pellet was resuspended again in rehydration solution, and the protein concentration was determined using the modified Bradford method.16 All IEF (Immobiline DryStrip gels [13 cm, pH 3–10]; GE Healthcare; rehydrated with 30 µg of solubilized proteins) and 2D electrophoresis steps were carried out as previously described.17 The proteins were resolved by electrophoresis through a 10% SDS-polyacrylamide gel (EttanTM DALT; GE Healthcare) at 5 watts (W)/gel for 30 minutes, followed by 17 W/gel for 4–5 hours. After electrophoresis, the gels were either silver stained or transferred onto a nitrocellulose membrane. Digital images were generated using transmission scanning (Image Scanner; GE Healthcare).

For 2D western blotting, Borrelia sp. proteins resolved by 2D gel electrophoresis (13 cm, pH 3-10) were transferred onto nitrocellulose membranes (Trans-blot Transfer Medium, pure nitrocellulose membrane, 0.45 µm; Bio-Rad, Ville-d’Avray, France). Membranes were then blocked in PBS supplemented with 0.2% Tween 20 and 5% non-fat dry milk (PBS-Tween-milk) for 1.5 hours at room temperature before incubation with MAbs P3A10 and P6D9 (dilution 1:100 or 1:500 for SDS-PAGE and western blot in the blocking buffer and a dilution of 1:1,000 for 2D western blotting). After 1 hour of incubation, membranes were washed three times with PBS-Tween and a dilution of 1:1,000 for 2D western blotting. After 1 hour of incubation, membranes were washed three times with PBS-Tween and probed with horseradish peroxidase-conjugated goat antihuman IgG (1:5,000; GE Healthcare). Membranes were subsequently incubated with the secondary antibody (biotin-conjugated antibody, 20 µg/mL) for 1 hour before three successive washes as described above. Immunostained spots were visualized using a commercially available chemiluminescence kit (ECL™ western blotting Analysis System; GE Healthcare). Then, the membranes were exposed to Hyperfilm™ ECL and subsequently developed using an automated film processor (Hyperprocessor™; GE Healthcare).

Digestion peptides and matrix-assisted laser desorption/ionization time-of-flight mass spectrometry analysis. The protein spots were excised from silver-stained gels either manually or by using Bio-Rad spot picker and identified using matrix-assisted laser desorption/ionization time-of-flight mass spectrometry analysis.
spectrometry (MALDI-TOF-MS) on a Bruker Ultraflex II spectrometer (Bruker Daltonics, Wissembourg, France) as previously described. After destaining, in-gel digestion with trypsin (sequencing-grade modified pore size trypsin; Promega, Madison, WI) was done as previously described. The peptides obtained from protein digestion were dissolved in 10−20 μL of 0.1% trifluoroacetic acid (TFA). The peptide mixture was then analyzed using MALDI-TOF-MS. The 0.3 μL peptide mixture was co-crystallized in the presence of 0.5% TFA onto the MALDI-TOF target with an equal amount of matrix solution (3 mg/mL of α-cyano-4-hydroxycinnamic acid in 50% acetonitrile). Alternatively, the peptide mixtures derived from protein digestion were desalted and concentrated using Zip tips (Millipore, Bedford, MA) and deposited onto the MALDI-TOF target by elution with the matrix solution. Mass spectra were internally calibrated using autolytic peptides from trypsin. The peptide mass fingerprints were used to identify the proteins, using Mascot software for comprehensive sequence databases. Searches were performed against all available sequences in public databases, including those for eukaryotes.

**Immunofluorescence detection of B. crocidurae.** Twelve O. sonrai were collected in Senegal, and the presence of B. crocidurae was confirmed in ticks as previously described. The ticks were tested for the presence of B. crocidurae by glpQ gene real-time polymerase chain reaction (PCR) using a Ct ≤ 35 cutoff. The blind immunofluorescence assay incorporating both antibodies P3A10 and P6D9 was done using hemolymph of ticks as described. The distal portion of a leg of tick was amputated, and the hemolymph that appeared at the leg extremity was smeared onto a microscope slide, stained by Giemsa staining or immuno-detection and fixed with methanol for 5 minutes. The slides were air-dried, and fixed with methanol for 5 minutes. The slides were saturated by incubation with PBS-5% secondary antibody FITC with a sample for secondary controls. For image scanning, slides were mounted with Fluoprep (bioMérieux) after subsequent washing procedures and examined under an Olympus BX-51 epifluorescence microscope (Olympus, Rungis, France).

Five 7-week-old female guinea pigs were inoculated intraperitoneally with 10⁷ B. crocidurae or buffer (one negative control). Blood obtained by cardiac puncture 5 days post-inoculation was tested by immunofluorescence assay incorporating P6D9 or P3A10 as described above.

Six blood smears from patients diagnosed by real-time PCR with B. crocidurae relapsing fever and from two patients negative for B. crocidurae in Senegal were analyzed by immunofluorescence assay incorporating P6D9 or P3A10.

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**RESULTS**

**Reactivity.** A total of 12 MAbs were produced and iso-typed as IgG1k, IgG2ak, IgG2bk, and IgM k (Table 2). Indirect immunofluorescence assay incorporating relapsing fever borreliae and 11 blood-borne organisms responsible for fever in West Africa showed that 11 MAbs reacted for relapsing fever borreliae, with MAb P5A7 also reacting with a Bartonella quintana clinical strain at 1:40 titer. Antibodies P3A10 and P6D9 exhibited the highest titers against the B. crocidurae Achema strain (Tables 1 and 2) and further reacted against the B. crocidurae 03-02 clinical strain in immunofluorescence assay (Figure 1). In silico, multispace sequence typing (MST) of B. crocidurae 03-02 strain found sequence type 6 (ST6), which differed in the five spacer sequences used from the reference strain Achema (ST13). In particular, P3A10 recognized the B. crocidurae Achema strain at a 1:500 dilution and the B. crocidurae 03-02 clinical strain at 1:1,000 dilution.

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**Table 2**

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MAbs = monoclonal antibodies; neg = negative; pos = positive.

Samples were considered positive when the fluorescence intensity of bacteria was comparable to that of polyvalent sera from immunized mice.
SDS-PAGE and western blotting of *B. crocidurae*. To identify the epitopes recognized by the MAbs, *B. crocidurae* Achema and *B. duttonii* Ly extracts were subjected to 2D electrophoresis and subsequent western blotting analysis. SDS-PAGE and western-immunobloting using P3A10 and P6D9 found no reactivity against *B. burgdorferi* while P6D9 recognized a variable-like protein (Vlp) (molecular weight 35.1 kDa in *B. crocidurae* and 35.2 kDa in *B. duttonii*) (Figure 2). P3A10 and P6D9 have been deposited in the Deutsche Sammlung von Mikroorganismen (DSM) collection (DSM ACC3255 and DSM ACC3256, respectively) as required by the Budapest Treaty on the international recognition of the deposit of microorganisms for the purposes of patent procedure.

**Immunofluorescence detection of *B. crocidurae***. Among the 12 *O. sonrai* collected in Senegal, four were positive for *B. crocidurae* DNA. Blind immunofluorescence assay incorporating P3A10 and P6D9 on the hemolymph, using the hemolymph test technique for detection of rickettsiae in ticks by immunofluorescence, found no *Borrelia* in non-infected ticks and *Borrelia*-like organisms in 2/4 infected ticks (Figure 4). No *Borrelia* was detected in the negative control animal whereas P3A10 and P6D9 detected *Borrelia*-like organisms in 4/5 challenged female guinea pigs inoculated intraperitoneally with *B. crocidurae* (Figure 4). Finally, immunofluorescence assay incorporating P6D9 or P3A10 detected *Borrelia*-like organisms in 6/6 blood smears from patients diagnosed by real-time PCR with *B. crocidurae* relapsing fever in Senegal and no organisms in two negative controls (Figure 5).

**DISCUSSION**

The direct detection of relapsing fever borreliae in clinical and vector specimens currently requires use of technically demanding, time- and resource-consuming molecular methods. Although the serological confirmation is based on either an immunofluorescence assay or enzyme-linked immunosorbent assay, this technique presents drawbacks because the antigenic variability of relapsing fever borreliae outer surface protein Vlp (proteins that, as a result of
FIGURE 3. Immunoreactive proteins with mouse monoclonal antibodies (MAbs) P6D9 and P3A10. *Borrelia crocidurae* and *Borrelia duttonii* proteins were resolved in the first dimension over a isoelectric point gradient of 3–10 followed by 12% linear sodium dodecyl sulfate polyacrylamide gel electrophoresis for the second dimension. Proteins were then detected by silver staining (*A* and *D* for *B. crocidurae* and *B. duttonii*, respectively) or transferred to a nitrocellulose membrane and probed with purified mouse MAbs P6D9 (*B* and *E*; and P3A10 (C) at 1:1,000 dilution. Standard molecular weight markers are indicated on the left in kilodaltons (kDa). Identified immunoreactive proteins, achieved by matrix-assisted laser desorption/ionization time-of-flight analysis, are indicated by protein name.

FIGURE 4. Immunofluorescence assay detection of *Borrelia crocidurae* in the *Ornithodoros sonrai* tick (arrow) (*A* for negative control and *B* for assay) and bloodstream of guinea pigs (*C* for negative control and *D* for assay) with P3A10 and P6D9 monoclonal antibodies (MAbs). Purified mouse MAbs P3A10 and P6D9 at 1:500 dilution were tagged using a goat anti-mouse IgG conjugated to fluorescein isothiocyanate at 1:400 dilution. Magnification: ×500 for guinea pig negative control and ×1,000 for other pictures.

FIGURE 5. Immunofluorescence assay detection of *Borrelia crocidurae* in blood smears from patients diagnosed with *B. crocidurae* relapsing fever in Senegal using P3A10 and P6D9 purified monoclonal antibodies. *A*, *B*, and *C* are blood smears of three patients; *D*, *E*, and *F* are negative controls. P3A10 and P6D9 at 1:500 dilution were tagged using a goat anti-mouse IgG conjugated to fluorescein isothiocyanate at 1:400 dilution. Magnification: ×1,000.
antigenic variation, allow relapsing fever borreliae to escape the host immune response and antigens shared with the Lyme disease spirochete (B. burgdorferi) may cause both false-negative and false-positive results, while serological tests based on the gp10 antigen can discriminate between relapsing fever and Lyme borreliosis. In this research based on the fact that B. crocidurae is the most common recognized Borrelia causing relapsing fever in West Africa, we produced 12 MAbs against B. crocidurae. Eleven of these MAbs reacted with borreliae and did not react with 11 blood-borne organisms responsible for fever in West Africa. The identification and characterization of the immunoreactivity of the two MAbs that exhibited the highest titers (P3A10 and P6D9) indicated that P6D9 cross-reacted with only B. crocidurae and B. duttonii Vlp. As Vlp is an antigen that varies in the surface protein of Borrelia, they may not be the stable and suitable targets for diagnostic MAbs. However, MAb P3A10 binds to flaB epitope 35.6 kDa, unique to B. crocidurae and may be a good candidate to be incorporated into a rapid diagnosis test for relapsing fever borreliae. Indeed, MAb P3A10 was shown to react equally with two cultured B. crocidurae isolates representative of two MST types. Furthermore, it detected B. crocidurae in six human blood smears in Senegal, two ticks and four animals, although these animals were experimentally infected and thus may not truly represent the sensitivity found during natural infection. We showed previously that several B. crocidurae genotypes were circulating in Senegal by MST, and these data suggest that MAb P3A10 recognized several genotypes in B. crocidurae. The identity of a Borrelia isolate can be classified according to protein profiles and immunoreactivities with specific MAbs. Although the diversity of major Osps and flagellar epitopes had been demonstrated in Lyme disease isolates from various geographical areas, for relapsing fever borreliae, P3A10 MAb recognized the flaB epitope of B. crocidurae. The most recent epidemiological data indicate that 43.92 million people living in rural Africa in endemic countries and 19.17 million travelers are at risk of relapsing fever in West and North African countries. Extrapolating on the 11% incidence of tick-borne relapsing fever measured in rural Senegal, this represents about 4.82 million cases of relapsing fever a year. Indeed, both antibodies could be incorporated into a format well suited for the rapid point-of-care diagnosis of relapsing fever in both endemic countries and in countries with travelers. In Africa, relapsing fever is often confused with malaria, therefore requires specific treatment and a specific prophylaxis.

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