Comparison of Enzyme-Linked Immunosorbent Assay, Western Blotting, Microagglutination, Indirect Immunofluorescence Assay, and Flow Cytometry for Serological Diagnosis of Tularemia

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The serodiagnostic efficiencies of five different approaches to detecting antibodies (immunoglobulins G, A, and M) developed in clinically proven infections with Francisella tularensis have been assessed. Fifty serum samples from patients suffering from tularemia during an outbreak in Sweden were compared with samples from 50 healthy blood donors (controls) by using an enzyme-linked immunosorbent assay (ELISA), microagglutination (MA), Western blotting (WB), an indirect immunofluorescence assay (IIFA), and flow cytometry (FC). ELISA, WB, and FC were based on the use of preparations of lipopolysaccharides (LPS) of the live vaccine strain of Francisella tularensis subsp. holarctica (ATCC 29684) as a capture antigen. Whole methanol-fixed bacteria were used for IIFA and MA. Optimized protocols yielded a diagnostic sensitivity and specificity of 100% for WB, MA, and FC, 98% for ELISA, and 93% for IIFA. A total of 6,632 serum samples from individuals between the ages of 18 and 79 years, representatively recruited from all regions of Germany, were screened to estimate and confirm the positive predictive value (PVpos) of the ELISA. Serum samples from 15 (0.226%) individuals tested positive for F. tularensis-specific antibodies by ELISA and confirmatory WB. The resulting prevalence-dependent PVpos of 10.2% and specificity of 98.1% were consistent with our findings for tularemia patients and controls. We conclude that the combined usage of a screening ELISA and a confirmatory WB based on LPS as a common antigen, as well as the MA, is a suitable serodiagnostic tool, while the quality of the IIFA is hampered by subjective variations of the results. FC is a promising new approach that might be improved further in terms of multiplex analyses or high-throughput applications.

Tularemia is a zoonotic disease caused by the highly infective, virulent, nonsporulating gram-negative coccobacillus Francisella tularensis. It is found throughout most of the northern hemisphere in a wide range of animal reservoir hosts including mammals, birds, and insects and can persist for a long time in contaminated environmental sources such as water and mud. It is not known to be transmitted from one person to another. Epidemics can often be traced to concurrent epizootics involving rodents and other small mammals (for a detailed review, see reference 15). Furthermore, in the past, tularemia was one of the most common laboratory-acquired diseases (10).

There are several tularemia syndromes in humans, most of them depending on the portal of infection. The clinical appearances range from skin lesions to multiorgan involvement. The severity, furthermore, varies with the dose inoculated and the virulence of the bacterium, which is related to the biotype. Francisella tularensis biovar holarctica (type B), which is spread over the whole northern hemisphere, is less virulent than Francisella tularensis biovar tularensis (type A), which exists mainly in North America and is associated with severe and often fatal tularemia if left untreated.

The usual incubation period is 3 to 5 days, although it can be as long as 21 days. In most cases, antibodies appear 6 to 10 days after the onset of symptoms, i.e., usually about 2 weeks after infection, reach their peaks at 4 to 7 weeks, and, despite decreasing in level, are still present 0.5 to 25 years later, probably even longer (16, 28). Even though early identification of the pathogen is important, neither isolation by cultivation, immunologic detection of antigens, nor molecular approaches are always successful or suitable (22, 25, 29, 35). Considering the facultatively intracellular localization of the pathogen, cell-mediated immunity is likely to be the best correlate for assessing exposure or immunity to F. tularensis (26). A tularin skin test is very helpful in this regard and is still used in some parts of Eastern Europe (12) but is not licensed in most other countries. Thus, clinical investigations and epidemiological studies on humans and animals or confirmation of immunoreactivity after vaccination depend on the availability of reliable, convenient, and affordable assays to detect and monitor the appearance of specific antibodies. During the immune response, epitopes of Francisella lipopolysaccharides (LPS) are the main target for the development of species-specific antibodies (1, 32). These antibodies, however, confer virtually no or low protection on mice against challenges with virulent type A strains of the pathogen (13). Several other antigens have been used to confirm seroconversion after exposure to F. tularensis (7). Preparations of outer membrane antigens can be applied to several methodological platforms such as enzyme-linked
immunosorbent assays (ELISA), microagglutination, and Western blotting (5, 6, 8). Assays based on LPS as a capture antigen, with different approaches to purification and detection, have been described previously (11, 20, 21, 37). These preparations have yielded high specificity for Francisella type A and B strains, the assays are easy to carry out, and antigens were stable over a long period (22).

In this study, we compared different preparations and platforms routinely used for the detection of F. tularensis-specific antibodies. The efficiencies of the different tests were assessed, and the plausibility of our findings was confirmed in a large-scale field investigation. Additionally, we set up a new approach toward further antigen purification and detection of specific antibodies, employing flow cytometry, a rapid detection technique which is increasingly used in clinical laboratories (36).

MATERIALS AND METHODS

Bacteria and LPS preparation. The live vaccine strain (LVS) of F. tularensis biovar holarctica (ATCC 29268) was grown over 2 days on heart-cytochrome blood agar and harvested into sterile distilled water, and bacterial concentrations were adjusted photometrically at 560 nm. Therefore, the LPS concentration is given as "bacterial equivalents." For the ELISA, the bacteria were adjusted to an optical density at 560 nm (OD560) of 1.0, which corresponded to 109 bacteria/ml. A 1/100 "bacterial equivalents. " For the ELISA, the bacteria were adjusted to an optical density at 560 nm. Therefore, the LPS concentration is given as the optimal concentration for coating the microtiter plates.

To extract and solubilize the LPS from bacteria, a Chlamydia specimen extraction buffer (Abbott, Wiesbaden, Germany) was added to a final dilution of 1/2 and incubated for 30 min at 60°C. It was not expected that a highly purified LPS preparation would be obtained by this procedure. However, in another set of experiments, we purified the LPS to a high degree by phenol-water extraction and compared the purified LPS with the standard product. We did not find an improvement in specificity due to the use of the highly purified product compared with the LPS preparation described here. It was therefore concluded that the purification described here is sufficient.

To further purify the LPS, the antigen was filtered and treated with 3.3 mg of proteinase K (Boehringer, Mannheim, Germany)/ml for 2 h at 60°C in order to digest residues of proteins. The enzyme was heat inactivated for 25 min. LPS was then precipitated by addition of 2.5 in PBS with 0.05% thiomersal and 0.1% crystal violet. This suspension of LPS was stable over a long period (22).

Microagglutination. Colonies of F. tularensis LVS were harvested into 0.9% NaCl from plates after cultivation for 2 days. After inactivation by 1% parafomaldehyde overnight, bacteria were washed twice and adjusted to an OD560 of 2.5 in PBS with 0.05% thiomersal and 0.1% crystal violet. This suspension of stained bacteria was used as an antigen. Serial dilutions of the sera up to 1/256 in PBS were applied. Fifty microliters of antigen suspensions was mixed with 50 µl of diluted sera by using 96-well round-bottom microtiter plates (NUNC), and the mixture was incubated overnight at room temperature. Wells were subsequently examined visually for a typically blue color indicating agglutination.

Flow cytometry. The IgG1 monoclonal antibody (MAb) 11/1/6 was produced by myeloma cells fused with mouse (BALB/c) spleen cells after immunization with F. tularensis ATCC 6225 as previously described (21). No cross-reactivity with a wide range of gram-negative bacteria was observed (22).

The assay is based on the coupling of MAb 11/1/6, specific for F. tularensis LPS, on magnetic M-450 goat anti-mouse-DynaBeads (Dynal, Hamburg, Germany) and the subsequent precipitation with F. tularensis LPS. The bead-antibody-LPS complex was incubated with sera. F. tularensis LPS-specific antibodies were detected by fluorescein isothiocyanate (FITC)-labeled goat anti-human IgG, IgA, and IgM antibodies (Sigma). Agglutination, centrifugation time, dilution of sera, antibodies, and antigen were pretreated in order to optimize assay settings. In the final protocol, 5 µl of beads (absolute number, 3 x 105) was intensively washed and resuspended in 1 ml of fluorescein-activated cell sorter (FACS) buffer containing PBS (pH 7.2; Sigma-Alrich, Taufkirchen, Germany), 2% fetal calf serum (Gibco), and 0.02% sodium azide (Merck, Darmstadt, Germany).

Single washing steps were usually performed. If intensive washing was required, the procedure was repeated three times. Beads were incubated with 40 µg of MAb 11/1/6/ml against F. tularensis LPS was 1/2 diluted and incubated at room temperature for 30 min with the beads. Loaded beads were then transferred to FACS tubes and incubated with the sera at a final dilution of 1/25 for 30 min. Polyvalent FITC-labeled anti-human immunoglobulin was diluted 1/100, and 200 µl was added to each tube. After incubation for 30 min at room temperature, samples were resuspended in 500 µl of FACS buffer and subjected to flow cytometry by using an EPICS XL (Beckman Coulter, Krefeld, Germany). Between incubation steps, beads were washed by magnetic separation using the MCP-1 magnetic particle collector (Dynal).

Indirect immunofluorescence assay (IFA). Suspensions of the LPS (OD560 1.0) were diluted 1/20, and 20 µl was placed on immunofluorescence slides (BioMerieux, Marcy l’Etoile, France). Slides were dried overnight and fixed with methanol. Serial dilutions of 20 µl of sera were added and incubated at 4°C for 1 h. FITC-labeled polyvalent anti-human immunoglobulins (Dako, Hamburg, Germany) were diluted 1/100 in PBS containing 0.05% Evans blue. The secondary antibodies used were found not to be reactive with Francisella antigen. After a wash with PBS, slides were covered with 20 µl of the secondary antibody-dye mixture, incubated at 37°C for 30 min, and washed with PBS. Slides
were subsequently covered with fluorescent mounting medium (DAKO). The fluorescence intensity was judged microscopically by the brightness, which corresponded to a semiquantitative scale from negative to quadruple-cross positive. Antibody titers were defined as the highest serum dilution that showed a specific fluorescence corresponding to double-cross positive (if not otherwise indicated). Lower brightness led to a relatively high subjective variation of the results when investigators were blinded.

**Statistics.** The diagnostic performance of a test was assessed by receiver operating characteristic (ROC) analysis including ROC curves, a complete sensitivity-specificity report, and disease prevalence-dependent predictive values (39). A sample size of 50 patients and 50 controls was used, according to the method of Metz (30), in order to draw meaningful conclusions from ROC experiments.

Linear regression and correlation analyses were performed in order to assess the relationship and association between two variables. MedCalc, version 7.2.0.2, was used for all statistical analysis purposes.

**RESULTS AND DISCUSSION**

Setup and comparison of serodiagnostic approaches. In the present study, five different methods of detecting *Francisella*-specific antibodies were evaluated. *F. tularensis* LPS is one of the major targets for antibodies (1, 32). While whole bacterial suspensions are needed for immunofluorescence tests and microagglutination, lysed bacteria were used for Western blotting. Further purification of LPS by elimination of contaminating proteins was necessary for ELISA and flow cytometry in order to reduce nonspecific binding of serum antibodies in presumptively nonexposed subjects. The diagnostic performance of these assays was evaluated by using random samples from 50 patients who had suffered from clinically proven tularemia and 50 healthy blood donors (controls). The distributions of antibody levels obtained by the different diagnostic approaches are depicted in Fig. 1. Calculations of sensitivity, specificity, and the underlying cutoff levels are given in Table 1 and are discussed below. A comparison of ELISA ODs at a serum dilution of 1/500 with end point titers of microagglutination and IIFA, respectively, showed no significant correlation (data not shown). It is most likely that the optical densities of positive and negative sera in ELISA were not measured in the linear range of the standard curve, so no conclusions can be drawn as to the end point titers. In contrast, mean fluorescence intensities obtained by flow cytometry from patients’ sera correlated significantly with the ODs of the ELISA (see below).

**Western blotting.** As shown in Table 1, Western blot analysis clearly distinguished between patients and controls. Dilution of sera down to 1/2,000 did not affect the appearance of the typical LPS banding pattern but eliminated nonspecific bands seen in some of the control sera. Cross-reactivity has been assumed for proteins homologous to the highly conserved chaperone proteins DnaK, GroEL, and GroES of *Escherichia coli* (17). Enzymatic digestion of the LPS extract decreased the amount of contaminating proteins. However, some heat shock proteins may have been resistant to proteinase K, since the

**FIG. 1.** Distribution of *F. tularensis* LPS-specific immunoglobulin levels. Levels in sera of patients (n = 50) and healthy controls (n = 50) were determined by the methods given above the plots. Sera for ELISA were diluted 1/640.
respective bands remained visible in the Western blot after digestion.

**IIFA.** Elevated optical densities in the screening ELISA and a high background immunofluorescence of negative controls might result from antibodies against those heat shock proteins. By IIFA, there was a clear overlap between patients and controls (Table 1). Decision criteria were defined as the highest titer of serum dilution that still allows the clear detection of fluorescence. Lower fluorescence intensities led to a higher subjective variation in determining the end point titers of sera.

**Microagglutination.** Microagglutination resulted in 100% sensitivity and specificity. No agglutination was visible in any of the control sera at dilutions below 1/16, while in 92% (n = 46) of the patients’ sera, titers reached 1/64 or higher (Fig. 1). Since the sera from all patients were collected more than 7 days after clinical manifestation of the infection, our results were in line with those of a former study that reported 24, 50, and 100% sensitivity of microagglutination 4, 5, and 7 days after clinical manifestation, respectively (33). These observations confirmed that microagglutination is a useful tool for the early and specific serodiagnosis of tularemia. However, cross-reactions with different species of *Brucella, Yersinia enterocolitica,* and *Proteus* must be considered and ruled out in the case of positive test results (2, 27).

**Flow cytometry.** The panel of routinely used methods was extended by an innovative approach that combines flow cytometry and immunomagnetic separation. Recently, flow cytometry has been reported to be a new platform for serological investigations of plague (36). We therefore set up a procedure to couple *F. tularensis* LPS on goat anti-mouse paramagnetic beads (GAM-Dynabeads; Dynal, Hamburg, Germany) and used MAb 11/1/6, specific for LPS of all *F. tularensis* subspecies but not for *Francisella novicida* (22), as a linker. The coupling of beads, MAb 11/1/6, and LPS formed stable complexes. The stability of the coated beads was tested by comparing freshly prepared beads with those coated at the beginning of the test period (7 days) and stored at 4°C. Measuring positive- and negative-control samples revealed no significant differences in

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**TABLE 1. Cutoff values, diagnostic sensitivity, and specificity of four approaches for the detection of tularemia-specific antibodies**

| Method                  | Unit                          | At 100% sensitivity | At 100% specificity |
|-------------------------|-------------------------------|---------------------|---------------------|
|                         | Cutoff                        | Specificity (%)     | (95% CI)            | Cutoff | Sensitivity (%) | (95% CI)                        |
| ELISA                   | Optical density               | 0.648               | 98.0 (89.3–99.7)    | 0.780  | 98.0 (89.3–99.7)|
| Flow cytometry          | Mean fluorescence intensity   | 1.59                | 100.0 (92.8–100)    | 1.59   | 100.0 (92.8–100)|
| Indirect immunofluorescence assay | Intensity, titer | 1/80                | 92.0 (80.7–97.7)    | 1/320  | 94.0 (83.4–98.7)|
| Microagglutination      | Agglutination, titer         | 1/16                | 100 (92.8–100)      | 1/16   | 100.0 (92.8–100)|
| Western blotting        | Visible LPS pattern, titer   | 1/2,000             | 100 (92.8–100)      | 1/2,000| 100.0 (92.8–100)|

* Fifty serum samples from clinically proven tularemia patients and 50 from healthy controls were analyzed for *F. tularensis* LPS-specific immunoglobulin. Methods were evaluated by ROC analysis. 95% CI, 95% confidence intervals.

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**FIG. 2. Determination of *F. tularensis* antibodies by flow cytometry.** Representative histograms of green fluorescence intensities of five serum samples from tularemia patients and two from healthy controls are shown.
mean fluorescence intensity between fresh and stored beads (data not shown). We did not test stability over a longer period, even though we assume that stability might last even longer. Since the usage of precoated beads saved about 2 h of assay time, beads were prepared on Monday and used over the week. As shown in the overlay graph (Fig. 2), the optimized assay settings allowed a sufficient distinction to be revealed between the fluorescence intensities of patients and controls. The lower background of controls in flow cytometry than in ELISA might be caused by the additional purification of the LPS antigen due to the binding and washing of antibody-bead complexes. Systematic analysis of patient and control sera revealed a complete distinction between the two groups (Fig. 1).

**ELISA.** Enzyme-linked immunosorbent assays are among the methods of choice for large-scale investigations in the context of outbreak scenarios or epidemiological surveillance studies (4, 6, 28). LPS preparations immobilized on Polysorb microtiter plates were identical to those used for flow cytometry. Nevertheless, flow cytometry was obviously superior to ELISA in terms of test efficiency. The scatter graph in Fig. 3 indicates that ELISA results for patient samples correlated significantly with mean fluorescence intensities obtained by flow cytometry. In contrast, ELISA results for the control sera spread over a wide range and did not correlate with the outcomes of corresponding measurements by flow cytometry. Dilution of sera down to 1/640 reduced the nonspecific background observed in control sera and increased the test efficiency of ELISA as assessed by ROC analysis. However, further dilution again led to an increased overlap of titers between patients and controls due to the decreased sensitivity of the assay (data not shown). A certain number of negative sera showed a relatively high background; however, these sera were probably overrepresented in this random sample. In larger studies (see “Seroepidemiology” below), the proportion of false-negative sera was much lower, even when a lower cutoff level was used.

Sensitivity and specificity assessed for optimized ELISA settings reached 98% and thus did not differ significantly from those reported from a former study that evaluated an ELISA with 57 tularemia cases (4). That assay was based on an outer membrane preparation that included several immunologic reactive antigens and reached a sensitivity of 93% for the simultaneous detection of IgG, IgA, and IgM and 97.5% for the separate analysis of immunoglobulin isotypes. Furthermore, Bevanger et al. described a competitive ELISA using a purified 43-kDa outer membrane protein that was tested with 23 tularemia patients (7). The sensitivity and specificity reported were 95.7 and 96%, respectively, after a serum dilution of 1/64.

**Seroepidemiology.** Assuming that the optimized LPS-based ELISA has a specificity of 98%, the pretest probability of the disease was calculated for a hypothetical range of disease prevalence. The prevalence-dependent positive predictive values of ELISA results are shown in Fig. 4.

Especially in a setting with a low serological prevalence of tularemia, such as seroepidemiological investigations conducted to estimate the risk of exposure or to survey for subclinical courses of tularemia, 95% false-positive results by
ELISA will appear, assuming there is a seroprevalence of 1:1,000.

In order to confirm our results, we tested 6,632 sera that were randomly collected in the context of a health survey initiated by the German health authorities (3). The cross-sectional sample was representative of the German population in the years 1997 to 1998. A cutoff level of 0.200 for performing confirmatory Western blotting was derived from measurement of the negative-control sera on 200 distinct microtiter plates and calculation of 3 times the standard deviation. We found 165 sera above the cutoff level. In contrast to the distribution of our initial ELISA results, shown in Fig. 1, the percentage of serologically negative individuals with high background levels was significantly lower. We therefore concluded that such sera were accidentally overrepresented in our initial sampling. Fifteen positive sera were confirmed by Western blotting (Fig. 5). Another eight sera had slightly elevated values but exhibited only a weak or incomplete LPS banding pattern. Since all sera confirmed as positive (Fig. 5, line b) were clearly separated from the cutoff value (Fig. 5, line a), we assumed 100% sensitivity of our approach. The calculated specificity of 98.1% resulted in a positive predictive value of 10.8% for the ELISA and matched our initial findings obtained by using tularemia patients as positive controls and healthy individuals as negative controls (Fig. 4). As a consequence, a seroprevalence of 226 per 100,000 was calculated and therefore has to be assumed for the German population.

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Several approaches have been adopted to assess the prevalence of ticks and animals infected with *F. tularensis* (9, 23, 38), and a serosurvey of landscapers in an area of tularemia endemcity has been conducted (18). According to the reports of these studies, some of which have been located in Europe, tularemia might make a significant contribution to tick-borne zoonoses in Germany, and the risk of exposure to carcasses or excrement of small mammals that carry the pathogen might be underestimated. Tick-borne tularemia appears to be a mild illness, characterized by fever and cervical or occipital lymphadenopathy (34). Persons at particular risk include hunters, as recently reported from Austria (14). Five out of 149 (3%) hunters examined exhibited antibodies against *F. tularensis*, without severe clinical symptoms. Besides tularemia, these individuals were also disproportionately seropositive for a wide variety of other zoonotic pathogens. Yet only sparse efforts have been made in Germany to conduct systematic surveil-

![FIG. 4. Tularemia prevalence-dependent predictive values of ELISA. Positive (filled circles) and negative (open circles) predictive values of ELISA for a hypothetical range of disease prevalence were calculated based on the results given in Table 1. Ninety-five percent confidence intervals are indicated.](http://cvl.asm.org/)
lance of natural foci of tularemia, such as hares (19), in order to forecast the enzootic situation and to take preventive measures. In addition, there is a need for increased awareness of the clinical entities that arise from such a zoonosis. Improved recognition of clinical syndromes, combined with laboratory investigations, will lead to higher diagnostic accuracy. Our study highlighted the availability of a variety of reliable laboratory means to ensure the diagnostic quality needed in this context.

Concluding remarks. Among the assays evaluated, the combined usage of ELISA and confirmatory Western blotting seems to be the most suitable approach for serodiagnosis of tularemia. Microagglutination has advantages in terms of costs and the equipment needed and is therefore preferable in laboratories without sophisticated equipment or during field investigations (31). However, cross-reactions with Brucella, Y. enterocolitica, and Proteus spp. have to be considered. Even though the preparation of the antigen for the indirect immunofluorescence assay is most convenient and implementation of the assay is simple, its employment is hampered by the lack of test efficiency. However, an experienced investigator can compensate for this disadvantage. Flow cytometry has been tested for the first time as a new platform for serodiagnosis of tularemia. Our initial findings suggest that in terms of test efficiency, it is superior to ELISA and comparable to Western blotting and microagglutination. The lack of commercial availability of diagnostics and the cost-intensive, highly sophisticated technical equipment required may hamper its usage in routine diagnostic work. However, the use of array-based applications might allow the development of high-throughput or multiplex analyses on this platform in the future.

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