Leptospirosis infections among hospital patients, Sarawak, Malaysia

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Abstract

Background: Leptospirosis diagnoses have increased in Sarawak, Malaysia in recent years.

Methods: To better understand the burden of disease and associated risk factors, we evaluated 147 patients presenting with clinical leptospirosis to local hospitals in Sarawak, Malaysia for the presence of Leptospira and associated antibodies. Sera and urine specimens collected during the acute illness phase were assessed via a commercially available rapid leptospirosis test (Leptorapide, Linnodee Ltd., Antrim, Northern Ireland), an ELISA IgM assay (Leptospira IgM ELISA, PanBio, Queensland, Australia) and a pan-Leptospira real-time PCR (qPCR) assay to estimate disease prevalence and diagnostic accuracy of each method. Microagglutination testing was performed on a subset of samples.

Results: Overall, 45 out of 147 patients (30.6%) showed evidence of leptospires through qPCR in either one or both sera (20 patients) or urine (33 patients), and an additional ten (6.8%) were considered positive through serological testing, for an overall prevalence of 37.4% within the study population. However, each diagnostic method individually yielded disparate prevalence estimates: rapid test 42.2% for sera and 30.5% for urine, ELISA 15.0% for sera, qPCR 13.8% for sera and 23.4% for urine. Molecular characterization of a subset of positive samples by conventional PCR identified the bacterial species as Leptospira interrogans in 4 specimens. A multivariate risk factor analysis for the outcome of leptospirosis identified having completed primary school (OR = 2.5; 95 CI% 1.0–6.4) and weekly clothes-washing in local rivers (OR = 10.6; 95 CI% 1.4–214.8) with increased likelihood of leptospirosis when compared with those who had not.

Conclusion: Overall, the data suggest a relatively high prevalence of leptospirosis in the study population. The low sensitivities of the rapid diagnostic test and ELISA assay against qPCR highlight a need for better screening tools.

Keywords: Leptospirosis, Sarawak, Malaysia, Diagnostics
Background

Leptospirosis has been on the rise in Malaysia in recent years [1, 2], where it has been designated a notifiable disease since 2010. The disease is biphasic in nature, with acute flu-like symptoms at the onset lasting for about a week, followed by antibody production and excretion of leptospires in the urine in the second week of illness. Mild cases may present with little to no symptoms, but severe infections can damage vital organs, trigger cytokine storms, and lead to death [3, 4]. In Malaysia, a peak of 7806 confirmed cases and 92 deaths was reported in 2014 [2], which suggests that leptospirosis is a significant public health threat locally [1, 2, 5], particularly considering that many cases may be under-reported or misdiagnosed [5, 6].

Leptospirosis is a zoonotic bacterial infection that causes widespread morbidity and disproportionately impacts people in regions with warm, humid climates in which the bacteria can easily proliferate [3, 6]. It is caused by a gram-negative spirochete of the genus *Leptospira*, which has 64 known genomic species and 300 identified serovars [7, 8]. Early identification and treatment of leptospirosis is challenging as symptoms of infection are non-specific and easily confused with other more common causes of acute febrile illness.

Given limited capacity for culture and molecular testing in many clinical settings, leptospirosis is commonly diagnosed by serology. Microscopic agglutination testing (MAT) is considered the reference test for leptospirosis serological diagnosis, but it has low sensitivity in the early stage of infection until measurable levels of antibodies develop in the blood [9]. Such delays in diagnosis may lead to missing the critical window for administering antibiotics. In addition, MAT requires a panel of live leptospires as well as highly-trained personnel, and only a few laboratories in Malaysia are able to offer this technique [5]. Alternatively, commercially available enzyme linked immunosorbent assays (ELISAs) can be used to detect antibodies in sera, with the added advantage that they can be standardized [10]. Studies have found some ELISA assays to be more sensitive than MAT at identifying IgM presence during leptospirosis infection [3, 6]. Early identification and treatment of leptospirosis is challenging as symptoms of infection are non-specific and easily confused with other more common causes of acute febrile illness.

When molecular testing is available, evidence of leptospires can be seen in sera samples before antibody levels become noticeably elevated [3, 10]. The bacteria can also be detected in cerebrospinal fluid and urine specimens as the infection progresses [3, 4, 11]; some studies have identified *Leptospira* excreted in urine samples within as little as seven days of symptom onset [11, 12]. Polymerase chain reaction (PCR) assays have been successful in detecting leptospires in both sera and urine, however, the specialized equipment and training required for this method make it impractical in lower-resourced laboratory settings where such assets are often not available.

Serological rapid diagnostic tests (RDTs) have been developed as user-friendly alternatives to these more rigorous laboratory diagnostics and are now commonly used to screen for suspected leptospirosis [10]. This is standard practice in Malaysia, where RDT-positive samples are further sent to a central laboratory for confirmatory MAT [2, 5, 7, 13, 14].

In this study, we sought to assess the diagnostic accuracy of commercially available RDT and ELISA assays against a newly developed pan-*Leptospira* real-time PCR molecular reference test [15] while estimating the prevalence of leptospirosis in Sarawak, Malaysia. We additionally tested the feasibility of using urine samples to diagnose suspected leptospirosis patients, and characterized disease presentation among the study population.

Methods

Study subject enrollment

Written consent was obtained from a parent and/or legal guardian of all participants under the age of 18. Children aged 7–18 were additionally asked to assent to study participation. Between February and September 2019, licensed medical officers at three primary hospitals in Sarawak, Malaysia identified patients that matched the U.S. Center for Disease Control and Prevention (CDC) case definition for clinical leptospirosis infection. Patients were invited to enroll in this cross-sectional study if they presented with either fever (axillary temperature measurement of more than 37.5°C by digital thermometer) or known exposure to *Leptospira* contaminated water, accompanied by at least two commonly associated symptoms of leptospirosis: myalgia, headache, jaundice, conjunctivitis, or rash [16]. Hospitals were located in both urban (Sibu Hospital) and rural settings (Kapit Hospital and Sarieki Hospital). Kapit, Sibu and Sarieki are located in a triangular point with intercity distances of 66 km to 222 km. Altogether, these three divisions cover 51,544.7 km² and account for 42.4% of the land area of Sarawak.

Once informed consent was obtained from the patient and/or their legal guardian, trained study staff collected a blood sample, and instructed the participant in midstream urine collection procedures. Participants were additionally asked to provide basic demographic, symptomatology, and exposure risk information through the completion of a brief questionnaire.

Sample processing

In the clinical laboratory, blood samples were left to coagulate for up to 60 min at room temperature before centrifugation at 2000×g for 10 min. Serum was...
Rapid diagnostic test

Initial RDT screening was performed by a laboratory technician at the enrolling hospital using a commercial latex agglutination test currently used for leptospirosis screening in Sarawak, Malaysia (Leptorapide, Linnodoe Ltd., Antrim, Northern Ireland). This test is designed to detect IgM antibodies for the rapid serological diagnosis of human leptospirosis [17]. This screening was done separately using sera and urine specimens. Even though this RDT was designed for use with serum, both serum and urine samples were tested using the same procedure as per manufacturer instructions. Briefly, 5 µl each of the Leptorapide reagent and specimen were applied to the provided agglutination card and then mixed. The agglutination card was rotated gently for 2–3 min and the positive/negative result was interpreted by visual comparison with the included score card. Specimens with ambiguous results were run a second time, and considered inconclusive if still unclear.

Microscopic agglutination test

In accordance with the Malaysian Ministry of Health protocol, acute sera samples with a positive RDT result were shared with the Institute for Medical Research in Kuala Lumpur, Malaysia for confirmatory testing. A positive MAT was defined as a single antibody titer ≥400, an equivocal MAT was defined as a single antibody titer between 1:50 and 1:200, and a negative MAT was defined as a single antibody titer < 1:50. In our study, convalescent sera samples were not available for MAT analysis.

ELISA assay

Once acute sera samples were received, technicians performed a Leptospira IgM ELISA assay (PanBio, Queensland, Australia) per the manufacturer’s instructions. Briefly, patient sera and controls were diluted 1:100 in diluent reagent then added to Leptospira antigen coated microwells (serovar: Patoc). The plate was incubated for 30 min at 37 °C. After washing the plate with PBS, 100 µl of horseradish peroxidase-conjugated anti-human IgM was added and incubated further for 30 min at 37 °C. After a second PBS wash, 100 µl of the tetramethylbenzidine substrate was added and the plate was incubated for 10 min at room temperature. The reaction was stopped with 100 µl of 1 M phosphoric acid. The absorbance value of each well was read at 450 nm wave length and interpreted in terms of Pan-Bio units, calculated using the absorbance of positive control serum, negative control serum, and cut-off of calibrators provided by the manufacturer. A Pan-Bio unit ≥11 was considered positive for the presence of Leptospira IgM antibodies.

qPCR

Total DNA was extracted from 140 µl each of the aliquoted sera and urine specimen according to the manufacturer’s instructions, using the QIAamp RNA Mini Kit (QIAGEN, Hilden, Germany) as recommended by El Bali, et al. [18]. The extracted DNA was then added to a master mix, containing 10 ul Sso Advanced Universal Probes Supremix (Bio-Rad, Hercules, California, USA), 4.4 µl ultra-pure water, and 0.2 µm each of 20 µM primers and probes targeting the 16S Leptospira gene, as described by Mohd Ali, et al. [15] in their Pan-Leptospira qPCR assay, specifically designed for the Malaysian context. The amplification protocol consisted of 95 °C for 3 min followed by 50 cycles of 95 °C for 30 s and 61.3 °C for 30 s. As the qPCR results were not used for clinical diagnosis, a conservative approach was used to consider specimens with threshold levels (Cq) below 40.0 positive for infection. Positive and negative controls were included in each run.

Molecular characterization

Extracted nucleic acid from specimen were shipped on dry ice to Duke University in North Carolina, USA for further characterization. At Duke, samples were screened by conventional PCR and, when positive (band at ~ 550 bp), Sanger sequencing (Eton Bioscience, Inc., Research Triangle, NC, USA) was done to determine the Leptospira genetic diversity at the specific level targeting the secY gene. Primers secY-F (5′-ATGCCGATCATT TTTGCTTC-3′) and secY-R (5′-CCGTCCTTAATT TAGACTTCTTC-3′) [19] were used to amplify 549-bp fragments. The PCR conditions consisted of a transcription step at 50 °C for 30 min and an initial denaturation
step at 95 °C for 15 min, followed by 45 cycles of denaturation at 94 °C for 30 s, annealing between 50 and 58 °C for 30 s, and extension at 72 °C for 1 min, with a final extension at 72 °C for 10 mins, see standard operating procedure in supplemental information.

**Statistical analyses**

Questionnaire data and laboratory results were entered into Microsoft Excel and then analyzed using SPSS Statistics version 21 (IBM, 2012) and R 3.5.1 (R Core Team, 2018), with epitools (Aragon, 2020), epiR (Stevenson, 2020), ggplot2 (Wickham, 2016) and caret (Kuhn, 2020) packages.

Participants were considered positive for leptospirosis if (i) either sera or urine showed evidence of Leptospira by qPCR, or (ii) at least two out of three serological tests (RDT, MAT, or ELISA) were positive. Sensitivity, specificity, and positive and negative predictive values (PPV, NPV) of the Leptorapide RDT and ELISA were calculated against both the combined and specimen-specific qPCR results. Correlation between tests was assessed using the phi coefficient.

Bivariate analysis using a student t-test and Fisher’s exact test were used to compare participant characteristics and their laboratory assays against the qPCR outcomes. Characteristics determined to be predictive of leptospirosis infection (p-value < 0.10) were incorporated into a generalized linear model, then further refined through stepwise, backward-elimination to identify statistically significant (p-value < 0.10) adjusted odds ratios.

**Results**

**Participant characteristics**

A total of 147 patients (88 male, 59 female) were enrolled across all three hospital sites (Table 1). Participant ages ranged from 2 to 78 years, with 56 participants (38.1%) under 12 years of age, 13 (8.8%) between 12 and 18 years, 71 (48.3%) between 18 and 65 years, and 7 (4.8%) above 65 years of age. Five different ethnicities were represented in the study, with the majority of participants at each location and in the overall study (122, 83.0%) identifying as Iban.

Many participants lived in longhouses (78, 53.1%), with 49 (33.3%) reporting residency in houses, and 20 (13.6%) in wooden huts. Households frequently reported obtaining drinking water from multiple sources; 65 (44.2%) collected water from the river, 72 (49.0%) had water piped into their homes, 17 (11.6%) used rainwater, and 10 (6.8%) had gravity feed water systems. River water was the most common drinking water source for participants enrolled at Kapit Hospital (55, 53.9%), while participants from Sarikei and Sibu Hospital more commonly reported using piped water (14 per site, 58.3 and 66.7%, respectively).

Among the study population, 34 (23.1%) had not completed any level of formal schooling, while 57 (38.8%) had completed primary and 49 (33.3%) completed secondary schooling. Only 7 (4.7%) participants had gone on to complete college or further professional studies. Among the 78 participants greater than 18 years of age, 46 (59.0%) were involved in occupations associated with high risk of leptospirosis infections, including agriculture (n = 29), fishing (n = 17), hunting (n = 11), and logging (n = 9). Many participants participated in multiple such activities. Within the study population, 52.6% of individuals greater than 18 years of age reported wearing closed-toe shoes for work (n = 41), while 30.8% wore open-toed shoes or sandals. Other reported personal protective equipment (PPE) usage included gloves (14.1%), eye protection (5.1%), and face masks (9.0%).

**Leptospirosis prevalence**

Serological RDT and ELISA results were available for all 147 patients, with calculated prevalences of 42.2% and 15.0% respectively. Out of 62 positive RDT sera specimens, 53 were sent for confirmatory MAT results, along with 5 inconclusive RDT samples and 1 negative. Regardless of RDT outcome, 15/59 (25.4%) sera samples returned positive MAT results. Per our positivity criteria (at least two out of three serological tests positive), a total of 23 participants (15.6%) were considered serology-positive for leptospirosis, of which thirteen participants also presented molecular evidence of leptospires in either sera or urine (Fig. 1).

Two sera samples did not have sufficient volume to complete qPCR testing. Among 145 remaining sera and 141 matched urine specimens, a total of 45 leptospirosis infections were identified through qPCR (positive for either sera and/or urine). Evidence of Leptospira was found in 20 (13.8%) sera and 33 (23.4%) urine specimens.

Taken together, serological and molecular testing identified 55 patients with leptospirosis, resulting in an estimated prevalence of 37.4% (55 out of 147) within our study population (Table 2). At Kapit Hospital, 34 of 102 patients (33.3%) were considered positive for leptospirosis, as compared to 9 of 24 (37.5%) at Sariekei Hospital, and 12 of 21 (57.1%) at Sibu Hospital.

**Diagnostic accuracy**

Leptorapide RDT analysis on patient sera samples identified 62 (42.2%) as positive, 67 (45.6%) as negative, and 18 (12.2%) inconclusive. RDT analysis on urine identified 43 specimens (30.5%) as positive and 98 (69.5%) as negative. Sensitivity, specificity, PPV, and NPV were assessed for each set of RDT analyses, using combined qPCR results for sera and urine as the gold standard (Table 3). MAT results were primarily available for RDT positive
samples, and thus excluded from the diagnostic accuracy calculations to avoid bias; of note, 45.8% of MAT results were inconclusive, as compared to 25.4% positive and 28.8% negative findings (Supplemental Fig. 1). Results of the Leptorapide RDT were further compared to specimen-specific qPCR results. The IgM ELISA assay identified 22 positive (15.0%), 113 negative (76.9%), and 12 equivocal specimens (8.2%). Sensitivity, specificity, PPV, and NPV of the IgM ELISA assay were calculated in comparison to combined qPCR results for sera and urine, as well as for sera-specific qPCR results (Table 4). As before, MAT results were excluded from this analysis.

**Molecular characterization**

SecY sequencing was performed on samples showing PCR amplification and a clean band of the expected size (549 bp) on gel electrophoresis. Samples successfully sequenced ($n = 4$) originated from urine specimen collected within five days of fever onset. After cleaning and trimming, we compared the 410-bp fragments to secY sequences available from the MLST database (https://...
Fig. 1 Comparison of positive serological leptospirosis results for patients presenting with febrile illness to study hospitals in Sarawak, Malaysia according to various diagnostic methods used. Participants were defined as having leptospirosis if at least two out of three serological tests were found to be positive and/or molecular evidence of *Leptospira* was identified through qPCR in either participant sera or urine. * RDT: Rapid diagnostic test, Leptorapide latex agglutination test (Linnodee, Ltd., Antrim, Northern Ireland). † MAT: Microagglutination testing, as carried out by the Institute of Medical Research reference lab in Kuala Lumpur, Malaysia, only partial results available. ‡ ELISA: *Leptospira* IgM ELISA assay (PanBio, Queensland, Australia).

Table 2 Leptospirosis prevalence estimates among study population (n = 147) at three enrolling hospitals, Sarawak, Malaysia, according to various diagnostic tests employed. Participants were defined as having leptospirosis if at least two out of three serological tests were found to be positive and/or molecular evidence of *Leptospira* was identified in either participant sera or urine.

| Diagnostic type   | Kapit Hospital n (%) | Sarikei Hospital n (%) | Sibu Hospital n (%) | Total n (%) |
|-------------------|----------------------|------------------------|---------------------|-------------|
| **Sera**          |                      |                        |                     |             |
| Leptorapide RDT*  | 35/102 (34.3)        | 12/24 (50.0)           | 15/21 (71.4)        | 62/147 (42.2) |
| MAT†              | 2/31 (6.5)           | 6/14 (42.3)            | 7/14 (50.0)         | 15/59 (25.4) |
| ELISA assay‡      | 14/102 (13.7)        | 3/24 (12.5)            | 5/21 (23.8)         | 22/147 (15.0) |
| qPCR              | 16/102 (15.7)        | 3/23 (13.0)            | 1/21 (4.8)          | 20/145 (13.8) |
| **Urine**         |                      |                        |                     |             |
| Leptorapide RDT*  | 32/100 (32.0)        | 4/22 (18.2)            | 7/19 (36.8)         | 43/141 (30.5) |
| qPCR              | 19/100 (19.0)        | 8/22 (36.4)            | 6/19 (31.6)         | 33/141 (23.4) |
| Leptospirosis-positive participants | 34/102 (33.3) | 9/24 (37.5) | 12/21 (57.1) | 55/147 (37.4) |

* Rapid diagnostic test: Leptorapide latex agglutination test (Linnodee, Ltd., Antrim, Northern Ireland). † Microagglutination testing, as carried out by the Institute for Medical Research reference lab in Kuala Lumpur, Malaysia, only partial results available. ‡ *Leptospira* IgM ELISA assay (PanBio, Queensland, Australia)
Table 3 Diagnostic accuracy of Leptorapide latex agglutination test (Linnodee, Ltd., Antrim, Northern Ireland) using acute sera and urine specimen, in comparison to real-time PCR (qPCR)

| Diagnostic Type | Serum-based RDT* | Urine-based RDT* |
|-----------------|------------------|------------------|
|                 | RDT vs qPCR on sera or urine (95% CI) | RDT vs qPCR on sera (95% CI) | RDT vs qPCR on urine (95% CI) |
| Apparent prevalence | 48.1 (39.2–57.0) | 30.5 (23.0–38.8) | 30.5 (23.0–38.8) |
| Sensitivity | 47.5 (31.5–63.9) | 22.2 (11.2–37.1) | 18.2 (7.0–35.5) |
| Specificity | 51.7 (40.8–62.4) | 65.6 (55.2–75.0) | 65.7 (56.0–74.6) |
| Positive predictive value | 30.6 (19.6–43.7) | 11.5 (4.7–22.2) | 14.0 (5.3–27.9) |
| Negative predictive value | 68.7 (56.2–79.4) | 81.8 (70.4–90.2) | 72.4 (62.5–81.0) |

* Rapid diagnostic test. † Calculated confidence interval

pubmlst.org/organisms/leptospira-spp) and identified *Leptospira interrogans* as the infecting species (GenBank accession numbers are provided in Supplemental Table 1). When compared to a worldwide database including 134 secY sequences of *Leptospira interrogans* [20], our four sequences were identical or closely related to leptospires identified in Malaysia, Indonesia, India, the Philippines and China.

Disease presentation

Among the 55 defined leptospirosis patients, 9.1% (n = 5) presented with severe disease, characterized by death (n = 1), septic shock (n = 4) and respiratory distress requiring invasive ventilation support (n = 3). Another seven (12.7%) presented with moderate disease requiring non-invasive ventilation support (n = 4), upper gastrointestinal bleeding (n = 1), atrial fibrillation (n = 1) and thrombocytopenia (n = 2). The remaining 43 patients (78.2%) presented with mild disease. Acute kidney injury and transaminitis were observed at all disease severity levels (34.5% and 32.7% respectively).

The most commonly self-reported symptoms among the leptospirosis-positive patients were fever (52, 94.5%), nausea or vomiting (39, 70.9%), headache (37, 67.3%), and chills (36, 65.5%) (Supplemental Table 2). Several patients reported experiencing myalgia (29, 52.7%), joint pain (26, 47.3%), or conjunctivitis (17, 30.9%), while lethargy and diarrhea were uncommon (3, 5.5% and 9, 16.4%, respectively). Notably, jaundice was not observed in any of the leptospirosis positive patients.

Of note, two additional study participants died during the course of their hospitalization, but we found no detectable evidence of *Leptospira* infection in either of their samples.

Symptom onset

Leptospirosis-positive patients generally presented in the early phase of the disease, indicating that they were likely in the acute stage of infection at the time of sample collection (Fig. 2). Overall, 47 of 55 defined positive patients (85.4%) were enrolled within 6 days post symptom onset (DPSO; self-reported), with five (9.1%) presenting on day 7 and one (1.8%) on day 8. Two patients did report having experienced joint pain and myalgia, among other symptoms, for 14 days.

Comparatively, patients considered negative for leptospirosis reported having experienced non-specific febrile symptoms from one to 28 days prior to sample collection.

Correlation calculations identified sera and urine speci men qPCR analysis as complementary (Supplemental Table 3). Of the 45 participants with molecular evidence for leptospirosis, 12 (26.7%) were qPCR positive only through sera analysis, 25 (55.6%) only through urine, and 8 (17.8%) had *Leptospira* positivity in both sample types (Table 5). Sera-based qPCR results primarily picked up evidence of *Leptospira* presence between 2

Table 4 Diagnostic accuracy of *Leptospira* IgM ELISA assay (PanBio, Queensland, Australia) using acute sera specimen, in comparison to real-time PCR (qPCR)

| Diagnostic Type | ELISA assay vs qPCR on sera or urine (95% CI) | ELISA assay vs qPCR on sera only (95% CI) |
|-----------------|---------------------------------------------|------------------------------------------|
| Apparent prevalence | 16.3 (10.5–23.6) | 16.4 (10.6–23.8) |
| Sensitivity | 24.5 (13.3–38.9) | 21.1 (6.1–45.6) |
| Specificity | 88.4 (79.7–94.3) | 84.3 (76.4–90.5) |
| Positive predictive value | 54.5 (32.2–75.6) | 18.2 (5.2–40.3) |
| Negative predictive value | 67.3 (57.8–75.8) | 86.6 (78.9–92.3) |
and 6 DPSO, with one patient testing positive with no documented fever (Fig. 3). Urine-based qPCR results were positive between 1 and 8 DPSO, with three patients testing positive with no documented fever (Fig. 4).

Participants with serological evidence for leptospirosis ($n = 23$) were identified from 1 to 8 DPSO, with one identified at 14 DPSO and one reporting no fever. Among these, the ten patients with no molecular evidence of *Leptospira* were primarily identified between 2 and 7 DPSO, with one presenting at 14 DPSO.

Independently of other serological assessments, the Leptorapide RDT provided positive results for patients 1 to 20 DPSO using sera samples and 1 to 14 DPSO using urine samples. The majority of the urine-based RDT positive results were within 1 to 7 DPSO, with 1 specimen positive at 10 DPSO and another at 14.

The IgM ELISA assay detected antibody evidence in patients 2 to 8 DPSO, and in one participant presenting without fever. MAT positives were identified primarily in participants 2 to 7 DPSO, as well as in one participant at day 10, one at day 14, and one presenting without fever.

**Risk factor analysis**

Bivariate analyses demonstrated the participant’s completed level of education, type of housing, and frequency of washing clothes in the local river as being associated with leptospirosis diagnosis. Participant ethnicity, occupation, use of PPE, primary water source, and exposure to cats, dogs, pigs, poultry, or rodents were not found to be significantly associated with leptospirosis diagnosis, nor was the enrollment site, recent flooding, engaging in

| Phase          | No. of Patients | DPSO* Range | Sera Mean Cq† (Range) | Urine Mean Cq† (Range) |
|----------------|----------------|-------------|-----------------------|------------------------|
| Onset          | 12             | 2–6         | 37.16 (35.36–39.45)   | –                      |
| Dissemination  | 8              | 0–5         | 36.57 (34.80–39.60)   | 35.77 (27.17–39.85)    |
| Excretion      | 25             | 0–8         | –                     | 37.38 (29.23–39.96)    |

*Days post-symptom onset (DPSO): fever duration at time of enrollment and sampling as self-reported by the patient. †Real-time PCR-determined cycle threshold (Cq)
other river-based activities, household income, or primary mode of transportation (Supplemental Table 4).

Multivariate analysis identified completing primary school and weekly washing of clothes in the local river as most predictive of a positive leptospirosis result in sera and/or urine. In this joint model, participants who had completed primary school were more than twice as likely to test positive for leptospirosis infection when compared to those with no education (OR 2.5, 95% CI 1.0–6.4). Those who completed secondary school were nominally less likely to show evidence of infection than those with no education (OR 0.9, 95% CI 0.4–2.5) (Table 6).

Washing clothes in the local rivers on a daily basis resulted in similar odds of having a leptospirosis diagnosis when compared to those who reported never washing clothes in the river (OR 0.9, 95% CI 0.3–2.3). In contrast, individuals who washed clothes weekly in the rivers had a tenfold increase in the likelihood of leptospirosis diagnosis, albeit with very low specificity (OR 10.6, 95% CI 1.4–214.8).

Multicollinearity across the model was not identified as a concern, with a mean variance inflation factor of 1.06 for all included variables.

**Discussion**

Our study data document the poor performance of the Leptorapide RDT assay compared to qPCR diagnosis. The screening tool displayed low sensitivity and specificity for both sera and urine specimen; over half the true cases would have gone unidentified through using only the RDT screening followed by MAT confirmation, as commonly prescribed in Malaysia. Insufficient data was available to provide effective comparison between the RDT and validated MAT results; the study of paired acute and convalescent sera samples would provide more conclusive MAT results and allow for better assessment of the antibody detection accuracy of the
Leptospira infections are often low when compared to PCR. When using qPCR as the gold standard test, the serum-based ELISA assay produced higher specificity than RDT, but much lower sensitivity. Results for the ELISA test may be improved by testing paired sera rather than only acute stage samples [23], how-ever this is more labor intensive and delays diagnosis.

There is a great need for more accurate, yet simple to perform diagnostic tests for leptospirosis in Southeast Asia (and other resource-restricted settings). PCR can confirm diagnosis in the early phase of the disease before antibody titers are at detectable levels, but appropriate laboratory support is lacking in many areas [24]. MAT testing is similarly resource intensive and restricted to reference laboratories, and misses a critical window as antibodies are lacking during the acute phase of the disease [5].

Our estimated prevalence data suggest that 37.4% (55 out of 147) of the patients who present with clinical signs and symptoms consistent with leptospirosis receive a positive laboratory confirmation. We recognize that this estimate has a number of limitations and may or may not represent true prevalence. However, our results support our position that leptospirosis is a frequent cause of illness in Sarawak, as reported in previous studies conducted in the region. From 2011 to 2013, Lela Suut et al. likewise found a population-based leptospirosis seroprevalence of 37.4% among 508 subjects among rural communities in the Rejang Basin of Sarawak, using ELISA and MAT [25]. They found evidence of three main serogroups detected in all three divisions: 22.1% Djasiman, 13.2% Shermani, and 7.9% Pomona [25]. A study conducted in urban areas of Sarawak describing the diversity of Leptospira spp. in rats and the environment identified four pathogenic species: L. interrogans, L. weilii, L. borgpetersenii and L. noguchii [26], while another study focusing on various rodent species in urban, suburban, and rural Sarawak identified L. interrogans and L. borgpetersenii [27]. In the present study, we did not perform serological typing but were able to characterize a subset of positive samples as infected by L. interrogans, a pathogenic species previously identified in human leptospirosis in the region [1, 28] and commonly reported worldwide.

To our knowledge, this is the first clinical study on human leptospirosis in Sarawak, East Malaysia. Most of the leptospirosis patients were enrolled during the first six days of fever onset, suggesting they were in the acute stage of the disease. Study data were remarkable in that no patients with laboratory evidence of leptospirosis infection exhibited jaundice, a finding that is inconsistent with previous reports from West Malaysia that reported higher prevalences (22.5% [29] and 4.9% [30]) of jaundice among their patients. As icteric leptospirosis is associated with higher mortality rates [3, 4], it is unsurprising that low patient mortality was attributed to acute infections in our patients, despite having been reported in other case series [29, 30].

Our data (Supplemental Fig. 2) support other studies that have also been able to detect Leptospira in urine during the acute phase (first week) of infection [11, 12]. Our study identified more urine samples as qPCR positive for leptospirosis than sera, which was unexpected as our samples were typically collected within a week post

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**Table 6** Predictive factors for positive leptospirosis diagnosis (having either molecular or serological evidence of leptospirosis) in the study population (n = 147), as calculated through bivariate and multivariate analyses

| Risk Factor | Positive cases (%) | Negative cases (%) | Unadjusted OR* (95% CI) | Adjusted OR* (95% CI) |
|-------------|--------------------|--------------------|-------------------------|-----------------------|
| Education completed |                   |                    |                         |                       |
| None (n = 34) | 10 (29.4)          | 24 (70.6)          | Ref‡                    | Ref‡                  |
| Primary (n = 57) | 28 (49.1)          | 29 (50.9)          | 2.28 (0.95–5.86)        | 2.47 (1.00–6.41)      |
| Secondary / College (n = 56) | 17 (30.4)          | 39 (69.6)          | 1.04 (0.41–2.74)        | 0.95 (0.36–2.53)      |
| Type of Housing |                   |                    |                         |                       |
| House (n = 49) | 23 (46.9)          | 26 (53.1)          | Ref‡                    | –                     |
| Hut / Wood house (n = 20) | 9 (45.0)          | 11 (55.0)          | 0.93 (0.32–2.67)        | –                     |
| Longhouse (n = 78) | 23 (29.5)          | 55 (70.5)          | 0.48 (0.22–1.00)        | –                     |
| Washing clothes in river |               |                    |                         |                       |
| Daily (n = 24) | 7 (29.2)           | 17 (70.8)          | 0.70 (0.25–1.78)        | 0.86 (0.30–2.27)      |
| Weekly (n = 5) | 4 (80.0)           | 1 (20.0)           | 6.03 (0.80–168.44)      | 10.56 (1.44–214.82)   |
| None (n = 118) | 44 (37.3)          | 74 (62.7)          | Ref‡                    | Ref‡                  |

* Odds ratio. † Calculated confidence interval. ‡ Reference category.
symptom onset. During the early stage of infection, *Leptospira* are most commonly investigated and identified within the bloodstream [3, 7]. Bacterial presence can later be seen in both sera and urine samples as the infection disseminates and *Leptospira* colonize the kidneys. Finally, leptospires are cleared from the bloodstream during the convalescent phase, while still present in urine. As described by Esteves, et al., different sample types may indeed prove more effective for identifying *Leptospira* infection at varying stages of disease [31], with the inclusion of urine permitting a wider window of detection [3]. By testing paired sera and urine specimens, we were able to identify 25 additional leptospirosis cases, which would have been missed had only sera been analyzed.

Risk factor data were interesting in that, while weekly washing of clothes in the river greatly increased disease risk when compared to those not using the river for such activities, doing so daily had minimal effect on this same risk. This finding was fairly imprecise, given the low number of observances of weekly clothes washing, however it may reflect longer exposure time in *Leptospira*-contaminated river water, if one assumes that once-weekly washing correlates with a larger amount of clothes to process. Despite taking place in the same body of water, other river-based activities such as swimming, bathing, fishing, or commuting were not significantly associated with disease risk within the study population, regardless of frequency.

Completing primary school appeared to greatly increase the risk of leptospirosis in comparison to those with either no formal education or with more advanced training. Although household income and known occupational risk factors for leptospirosis were not found to significantly impact leptospirosis outcomes in our study population, the increased risk associated with having completed primary school may serve as a proxy for other socioeconomic exposures not studied here. Though the Malaysian Ministry of Health has reported greater rates of leptospirosis within 25–60 year old individuals [2], age was surprisingly not found to increase disease risk in our bivariate analysis and thus not incorporated into our multivariate modeling.

The study had a number of limitations. First it was a short-term study with limited geographical spread; we sampled for only a short time and only at three hospitals among the 22 hospitals in Sarawak so our findings may not be representative of the entire state of Sarawak.

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**Fig. 5** There are multiple potential sources of *Leptospira* infections in the region. Much of the area around Kapit Hospital has dense tropical forest and riverine systems rich in wildlife. This wildlife may introduce *Leptospira* into water systems. Most people live in longhouses or farm huts on the outskirts of the town of Kapit. Transportation is chiefly via boats on rivers. While Kapit is supplied with treated water, the source of the water in rural areas is either from river water or gravity fed rain-water-systems. Images here illustrate the living and transportation environments in which this study took place (photos provided by Dr. King-Ching Hii).
Further, in accordance with local clinical diagnostic procedures, MAT assays were only performed on a subset of the acute sera specimens that had positive RDT findings. Hence, our prevalence estimates may have underestimated true prevalence. Additionally, occupational exposure was difficult to assess, as only half the participants were of working age and few self-reported engaging with previously recognized high-risk occupations (e.g., fishing and agriculture). As people from rural Sarawak have many environmental water exposures which could serve as the source of Leptospiira infection (Fig. 5), it is challenging to implicate specific risks and plans to study particular interventions. Finally, risk factor and symptom data were self-reported and could not be verified, offering the possibility of bias.

Conclusion
This study confirms leptospirosis endemicity among the study population in Sarawak, Malaysia and identified low diagnostic performance of the commercial rapid diagnostic test and IgM ELISA assay studied when used during the acute phase of the disease. We conclude that there is a critical need for more sensitive, inexpensive, and easy to use rapid diagnostics for leptospirosis in this region. Until such diagnostics are available, clinicians in areas where molecular testing is unavailable will be wise to continue to treat empirically if the clinical suspicion of leptospirosis is high and RDTs are negative. Finally, our study found that in areas where PCR-based diagnostics are possible, testing both sera and urine can increase the probability of detecting leptospirosis during the acute phase.

Abbreviations
CDC: U.S. Center for Disease Control and Prevention; IgG: Immunoglobulin G; IgM: Immunoglobulin M; ELISA: Enzyme-linked immunosorbent assay; MAT: Microscopic agglutination testing; NPV: Negative predictive value; PBS: Phosphate-buffered saline solution; PCR: polymerase chain reaction; PPV: positive predictive value; RDTs: Serological rapid diagnostic tests; rPCR: real-time polymerase chain reaction

Supplementary Information
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Additional file 1.

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Authors’ contributions
KH. and E.R. wrote the manuscript with support from V.G., M.G., R.B., and G.G. All authors contributed to editing and review of the manuscript. I.S., A.B., and J.G. carried out primary laboratory analyses. N. A and L.X. validated laboratory findings and facilitated sequencing. E.R. performed principle statistical analysis in association with V.G. and J.M. KH, D.G., and G.G. conceptualized and designed the study. KH. and G.G. supervised the project.

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Availability of data and materials
The data collected during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
The study was conducted in accordance with the ethical standards of the Helsinki Declaration and received a scientific review by and ethical approval from the Medical Research and Ethics Committee of the Ministry of Health, Malaysia (NMMR-18-1022-40334) and Duke University Institutional Review Board (Pro00100246), with extramural research review in accordance with the U.S. Navy Human Research Protection Program (HPR0.NAMRU2.2019.00003), in compliance with all applicable federal regulations governing the protection of human subjects.

Consent for publication
Not applicable.

Competing interests
The authors have no financial interests to disclose. Dr. Gray serves on editorial board for Tropical Diseases, Travel Medicine and Vaccines.

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References

1. Garba B, Bahamar AN, Khanani-Bejo S, Zakaria Z, Mutalib AR. Retrospective study of leptospirosis screening in Malaysia. Ecohealth. 2017;14(2):389–98. https://doi.org/10.1007/s10393-017-1234-0.

2. Zainudin A, editor Epidemiology and Current Situation of Leptospirosis in Malaysia. Perdana Kephatan Persekutuan Pihak Berkasuan Tempatan; 2015 September 8–9, 2015: Labuan: Ministry of Health Malaysia.

3. Levert PN. Leptospirosis. Clin Microbiol Rev. 2001;14(2):296–326. https://doi.org/10.1128/CMR.14.2.296-326.2001.

4. Haake DA, Levett PN. Leptospirosis in humans. Curr Top Microbiol Immunol. 2013;378:65–97. https://doi.org/10.1007/978-3-662-45059-8_5.

5. Garba B, Bahamar AN, Bejo SK, Zakaria Z, Mutalib AR, Bande F. Major epidemiological factors associated with leptospirosis in Malaysia. Acta Trop. 2018;178:242–7. https://doi.org/10.1016/j.actatropica.2017.12.010.

6. Costa F, Hagan JE, Calcagno J, Kane M, Torgerson P, Martinez-Silveira MS, et al. Global morbidity and mortality of leptospirosis: a systematic review. PLoS Negl Trop Dis. 2015;9(10):e0003898. https://doi.org/10.1371/journal.pntd.0003898.

7. Picardeau M. Diagnosis and epidemiology of leptospirosis. Med Mal Infect. 2010;40(9):421–9. https://doi.org/10.1016/j.medmi.2010.07.002.

8. Vincent AT, Schiettekatte O, Goarant C, Neela VK, Bernet E, Thibeaux R, et al. Serological prevalence of leptospirosis among rural communities in the Rejang Basin, Sarawak, Malaysia. Asia Pac J Public Health. 2016;28(5):450–7. https://doi.org/10.1177/1010539516648003.

9. Pui CF, Billung LM, Apun K, Su‘ut L. Diversity of Leptospira spp. in Rats and Environment from Urban Areas of Sarawak, Malaysia. J Trop Med. 2017;2017:3760674.

10. Blasdell KR, Morand S, Perera D, Firth C. Association of rodent-borne Leptospira spp. with urban environments in Malaysian Borneo. PLoS Negl Trop Dis. 2019;13(9):e0007720. https://doi.org/10.1371/journal.pntd.0007720.

11. Courdurie C, Le Govic Y, Bourhy P, Alexer D, Pailla K, Theodose R, et al. Evaluation of different serological assays for early diagnosis of leptospirosis in Martinique (French West Indies). PLoS Negl Trop Dis. 2017;11(6):e0005678. https://doi.org/10.1371/journal.pntd.0005678.

12. WHO. Human leptospirosis: guidance for diagnosis, surveillance and control. World Health Organization. 2003.

13. Bal AE, Gravekamp C, Hartskeerl RA, De Meza-Brewster J, Korver H, Terpstra HJ. Detection of leptospires in urine by PCR for early diagnosis of leptospirosis. J Clin Microbiol. 1994;32(8):1894–8. https://doi.org/10.1128/jcm.32.8.1894-1898.1994.

14. Bahaauud O, Pastuszka A, Le Bruin C, Ehrmann S, Lanotte P. Seven Years Leptospirosis Follow-Up in a Critical Care Unit of a French Metropolitan Hospital: Role of Real Time PCR for a Quick and Acute Diagnosis J Clin Med. 2020;9, 9, DOI: https://doi.org/10.3390/jcm9093011.

15. Bajani MD, Ashford DA, Bragg SL, Woods CW, Aye T, Spiegel RA, et al. A prospective hospital study to evaluate the diagnostic accuracy of rapid diagnostic tests for the early detection of leptospirosis in Laos. Am J Trop Med Hyg. 2018;98(4):1056–60. https://doi.org/10.4269/ajtmh.17-0702.

16. Desakorn V, Wuthiekanun V, Thanachartwet V, Sahasasanada D, Chierakul W, Apiwattanaporn A, et al. Accuracy of a commercial IgM ELISA for the diagnosis of human leptospirosis in Thailand. Am J Trop Med Hyg. 2012;86(5):524–7. https://doi.org/10.4269/ajtmh.2012.11-0423.

17. Nalam K, Ahmed A, Devi SM, Francalaci P, Baig M, Sechi LA, et al. Genetic affinities within a large global collection of pathogenic Leptospira: implications for strain identification and molecular epidemiology. PLoS One. 2010;5(8):e12637. https://doi.org/10.1371/journal.pone.0012637.

18. Chang CH, Riaz M, Yunus MH, Osman S, Noordin R. Limited diagnostic value of two commercial rapid tests for acute leptospirosis detection in Malaysia. Diagn Microbiol Infect Dis. 2014;80(4):278–81. https://doi.org/10.1016/j.diagmicrobio.201408012.

19. Dittrich S, Bouthraaoung L, Keshamharhong D, Phuklia W, Craig SB, Tulsiani SM, et al. A prospective hospital study to evaluate the diagnostic accuracy of four rapid diagnostic tests for the early detection of leptospirosis in Laos. Am J Trop Med Hyg. 2018;98(4):1056–60. https://doi.org/10.4269/ajtmh.17-0702.

20. Nalam K, Ahmed A, Devi SM, Francalaci P, Baig M, Sechi LA, et al. Genetic affinities within a large global collection of pathogenic Leptospira: implications for strain identification and molecular epidemiology. PLoS One. 2010;5(8):e12637. https://doi.org/10.1371/journal.pone.0012637.