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Human Leptospirosis Infection in Fiji: An Eco-epidemiological Approach to Identifying Risk Factors and Environmental Drivers for Transmission

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Abstract

Leptospirosis is an important zoonotic disease in the Pacific Islands. In Fiji, two successive cyclones and severe flooding in 2012 resulted in outbreaks with 576 reported cases and 7% case-fatality. We conducted a cross-sectional seroprevalence study and used an eco-epidemiological approach to characterize risk factors and drivers for human leptospirosis infection in Fiji, and aimed to provide an evidence base for improving the effectiveness of public health mitigation and intervention strategies. Antibodies indicative of previous or recent infection were found in 19.4% of 2152 participants (81 communities on the 3 main islands). Questionnaires and geographic information systems data were used to assess variables related to demographics, individual behaviour, contact with animals, socioeconomics, living conditions, land use, and the natural environment. On multivariable logistic regression analysis, variables associated with the presence of Leptospira antibodies included male gender (OR 1.55), iTaukei ethnicity (OR 3.51), living in villages (OR 1.64), lack of treated water at home (OR 1.52), working outdoors (1.64), living in rural areas (OR 1.43), high poverty rate (OR 1.74), living <100m from a major river (OR 1.41), pigs in the community (OR 1.54), high cattle density in the district (OR 1.04 per head/sqkm), and high maximum rainfall in the wettest month (OR 1.003 per mm). Risk factors and drivers for human leptospirosis infection in Fiji are complex and multifactorial, with environmental factors playing crucial roles. With global climate change, severe weather events and flooding are expected to intensify in the South Pacific. Population growth could also lead to more intensive livestock farming; and urbanization in developing countries is often associated with urban and peri-urban slums where diseases of poverty proliferate. Climate change, flooding, population growth,
urbanization, poverty and agricultural intensification are important drivers of zoonotic disease transmission; these factors may independently, or potentially synergistically, lead to enhanced leptospirosis transmission in Fiji and other similar settings.

Author Summary

Leptospirosis is a bacterial infection transmitted from animals to humans, and many outbreaks are associated with flooding. Globally, leptospirosis is responsible for at least a million cases of severe illness each year, and many deaths. The bacteria are excreted in the urine of infected animals; humans can become infected through direct contact with animals or through contaminated water and soil. In Fiji, two successive cyclones and severe flooding in 2012 resulted in 576 cases and 40 deaths. We conducted this study to improve our understanding of the factors that increase the risk of leptospirosis transmission, so that public health control measures can be improved. Our study found that infection risk is related to many factors including individual demographics and behaviour, contact with animals, living conditions, poverty, and flooding risk. With global climate change, flooding is expected to become a bigger problem in the South Pacific. Population growth could lead to more intensive livestock farming; and urbanization in developing countries is often associated with slums with high risk of infectious diseases. Climate change, flooding, population growth, urbanization, poverty and livestock farming are important factors for leptospirosis transmission; these factors may combine to increase the risk of leptospirosis in Fiji and other Pacific Islands in the future.

Introduction

Leptospirosis is an emerging infectious disease worldwide, with particularly high incidence reported in the Pacific Islands [1,2]. Humans are infected through direct contact with infected animals, or through contact with water or soil that has been contaminated by urine of infected animals. Disease transmission is strongly driven by environmental factors including high rainfall, flooding, natural disasters, population growth, urbanisation, and poor sanitation and hygiene [2–4]. In addition, infection risk depends on individual behaviour (e.g. swimming in fresh water, working outdoors), and contact with animals including livestock, rodents, pets, & wildlife [2,4]. Risk factors for infections and drivers of outbreaks depend on interactions between humans, animals, and the environment, and vary significantly between locations based on environmental, cultural, and socio-demographic factors [4]. Transmission dynamics are therefore highly complex and variable, and likely to evolve with global environmental change of both natural and anthropogenic environments [2,3].

In Pacific island nations, important risk factors for human leptospirosis include outdoor activities, tropical climate, flooding secondary to extreme weather events, and exposure to livestock [5–8]. Subsistence livestock are commonly kept in backyards, and veterinary expertise is generally limited. In some Pacific Islands, rapid population growth and urbanization exacerbate problems with sanitation, access to clean water, and waste management. Most islands have limited human or financial resources for the management and mitigation of the health impacts of natural disasters and climate change [9,10]. In Fiji, leptospirosis was identified as one of the four priority climate-sensitive diseases of major public health concern [11]. A recent systematic review of the global morbidity and mortality of leptospirosis identified tropical
islands as particularly high-risk settings [2]. Apart from the tropical climate and high frequency of extreme weather events [3], factors that could contribute to the high risk of leptospirosis on tropical islands include the low biodiversity and delicate ecosystems that make islands vulnerable to invasive species such as rodents [12]; the outdoor lifestyle and associated intense exposure to the environment; and close contact with subsistence livestock animals [2,4].

Climate change is projected to increase the severity of extreme weather events including increased rainfall and flooding in the Pacific Islands [10], and such events have been associated with increased leptospirosis transmission and outbreaks around the world [3,7,13–16]. In 2012, two successive tropical depressions caused severe flooding and resulted in two outbreaks of leptospirosis in Fiji, with 576 reported cases and 40 deaths (7% case-fatality) (Fiji Ministry of Health and Medical Services [MHMS]). Cases were defined as positive reactions to Leptospira ELISA IgM (Panbio, Brisbane, Australia); this laboratory test was only available at the national reference laboratory, and it was likely that reported cases were an underestimate of the true scale of the outbreaks. In comparison, previous studies in Fiji reported a total of 576 cases during an 8-year period from 2000–2007 [17], and 487 cases during a 13-year period from 1969–1981 [18]. These studies identified a higher risk of infection in males, indigenous Fijians (iTaukei), young adults (aged 15 to 45 years), rural dwellers and abattoir workers; increase in reported cases in the rainy months and after a cyclone in 2001; and geographic variation in incidence.

 Following the outbreaks in 2012, the Fiji MHMS and the World Health Organization convened a leptospirosis expert consultation to review the epidemiology of leptospirosis in Fiji and recommend priorities for control of endemic and epidemic disease. A key conclusion of the expert consultation was that significant knowledge gaps in the current epidemiology of leptospirosis in Fiji limited effective prevention and control. The study described in this paper was identified as one of several important steps to address the knowledge gaps. This study uses an eco-epidemiological approach and framework [19] to characterize the epidemiology and risk factors for human leptospirosis infection in Fiji, and aimed to provide an evidence base for improving the effectiveness and efficiency of public health mitigation and intervention strategies. Our findings would also be relevant to other countries with similar environments, particularly in the South Pacific.

**Methods**

**Study location and population**

The Republic of the Fiji Islands is an archipelago of 322 islands with a population of 837,217 in 2007; indigenous Fijians (iTaukei) and Indo-Fijians (Fijians of Indian descent) account for 57% and 35% of the population respectively [20]. Fiji is considered a ‘small island developing state’ by the United Nations [21] with a per capita GDP of US$4,712 [22]. The main island of Viti Levu has a landmass of 10,349 square kilometers and is home to >70% of the population. Vanua Levu is the second largest island in both population and land area, followed by Taveuni. The largest urban centre is the Greater Suva Area (population ~180,000) on the southeast coast of Viti Levu. The largest administrative units in geographical size are Divisions (Central, Western, Northern, and Eastern) followed by Provinces (14 in total), Tikinas (86 in total), and Enumeration Areas (smallest unit for population census that typically include 80 to 120 households). Nursing zones are the smallest administrative unit of the MHMS; they are under the care of a single nursing station and form a contiguous network across the Fijian Islands. Communities are residential clusters used by MHMS for administrative purposes. The four main community types in Fiji are urban residential areas, villages, Indo-Fijian settlements, and mixed Indo-Fijian/iTaukei settlements.
Seroprevalence study and sampling design

Field data were collected from September to December 2013 (January to March being the wettest months), and included the Central Division (on the eastern side of Viti Levu), the Western Division (on the western side of Viti Levu), and the Northern Division (the islands of Vanua Levu and Taveuni). The Eastern Division, with a population of ~40,000 spread across multiple small islands groups, was not included in the study because of logistical reasons. Field data were collected for a sero-epidemiological study of typhoid as well as the leptospirosis study described here.

We conducted a cross-sectional seroprevalence study, with a four-stage sampling design. An overview of the sampling plan is shown in Fig 1. In the first stage, both population-proportionate sampling and purposeful sampling approaches were used. The former was used to select 28 nursing zones from the Central Division, 21 from the Western Division, 11 from the Northern Division and 4 from the Ba Province which lies within the Western Division. Due to high incidence of reported leptospirosis and post-flood outbreaks in 2012, the latter sampling

![Fig 1. Overview of sampling strategy used in 2015 field study.](doi:10.1371/journal.pntd.0004405.g001)
approach was used to select 6 more nursing zones from the Ba Province. Similar to Ba Province, Taveuni Island (part of the Northern Division) was oversampled because of a high incidence of typhoid in 2008–2009. Consequently, 12 additional nursing zones were selected from this region, resulting in 82 zones in total being selected from the five regions in the first stage of sampling. At the second stage of sampling, one community was randomly selected from each of the 82 nursing zones. Headmen, health workers and other community leaders were consulted to seek agreement to participate in the study; no community leaders declined participation. At the third stage of sampling, 25 households were randomly selected from each community using health census records if available, or using a modified World Health Organization’s Expanded Programme on Immunization (EPI) sampling method. For the fourth and final stage of sampling, household members (defined as a person who stayed at the house the previous night) were enumerated and one selected at random for inclusion. In Ba subdivision, up to three randomly selected household members were included. If a selected household member was absent but returning later that day, the survey team would await their return or made a repeat visit. Wholly absent household members were substituted from within the household. Empty households were substituted by selecting the nearest house to the right of the front door. The above sampling strategy aimed to include 25 households from each of 82 communities, with up to three participants per household in Ba, and one participant per households in other areas. We therefore aimed to recruit a total of 2050 to 2250 participants.

Participants were eligible for inclusion if they were aged 12 months or older. Exclusion criteria included clotting disorders or medical anticoagulation, severe underlying medical conditions, significant acute illness, and fear of needles.

The communities included in the study represented the general population and different environments in Fiji (urban, peri-urban, rural), with higher sampling density in Ba and Taveuni. The study successfully included a total of 81 communities, with 28 in Central Division, 10 in Ba, 21 in other parts of the Western Division, 11 in Taveuni, and 11 in Vanua Levu. These areas will be referred to as the five ‘regions’ in this paper.

**Informed consent and ethics approvals**

Ethics approvals were granted by the Fiji National Research Ethics Review Committee (2013 03), the Human Research Ethics Committee of The University of Queensland (2014000008) and the London School of Hygiene & Tropical Medicine (6344). Support was sought and obtained from divisional and sub-divisional Ministry of Health officers for community visits. To ensure that research activities were culturally acceptable and local customs respected, community visits were conducted with field teams that included multilingual local Fijians. The study was explained to the heads of each of the randomly selected households, or another competent adult, and permission sought to include the household in the study. Written or thumbprinted informed consent was obtained from adult participants. The ethics committees specifically approved the use of thumbprint informed consent in illiterate participants. Parental/guardian consent and informed assent was obtained for child participants.

An overview of data sources and statistical methods is shown in Fig 2.

**Data collection during field study**

The following were collected from each participant:

1. Venous blood samples, collected by trained phlebotomists under sterile conditions (5–8mL depending on the age of the participant).
2. Questionnaire data, using standard questionnaires administered by field research assistants, and conducted in English or other local languages depending on each participant’s preference. Questions related to demographics, income, occupation, recreational activities, household environment, contact with animals, and other potential risk factors for leptospirosis.


Environmental, census, socio-demographic and livestock data

Environmental data on hydrology and roads were obtained from the Fiji Ministry of Lands and Mineral Resources [23]; and soils and land use/cover data from Fiji Ministry of Agriculture [24]. Climate (temperature and rainfall) and elevation data were obtained from the Landcare Research Institute [25]. Data on educational attainment, household construction, employment, ethnicity, and other socio-demographic variables were obtained from the 2007 Fiji National
Livestock data were provided by the Fiji Ministry of Agriculture’s 2009 National Agricultural Census [27]. All geospatial data were georeferenced to the Fiji Map Grid 1986 coordinate system. Table 1 provides a summary of the environmental, census, socio-demographic and livestock data used in the study. The five geographic regions used in this study and examples of the geo-referenced data are shown in Fig 3.

Household GPS coordinates from the study were projected on to the Fiji Map Grid 1986 coordinate system. Attributes from the geospatial predictor layers were linked to each household location by intersecting points through polygons for vector GIS data, and sampling the raster GIS data in a similar fashion. As a result, GIS attributes for each predictor layer were obtained for each household location. Attributes for a community were obtained by first calculating the location of the median centre of sampled households, followed by an approach similar to that which was carried out for individual households. All GIS data preparation and analysis was performed using ArcGIS v10.1 (Environmental Systems Research Institute, Redlands, CA).

All GIS data preparation and analysis was performed using ArcGIS v10.1 (Environmental Systems Research Institute, Redlands, CA).

Stratification of independent variables

Independent variables were stratified according to the ecological level at which they could potentially influence the risk of leptospirosis transmission and infection. Individual-level data relate to risk factors that are specific to individual demographics or behaviour. Household-level data

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Table 1. Summary of environmental, census, socio-demographic and livestock data used.

<table>
<thead>
<tr>
<th>Data</th>
<th>Source</th>
<th>Variables examined</th>
<th>Description &amp; Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrology</td>
<td>Fiji Ministry of Lands and Mineral Resources. Digital data from FLIS</td>
<td>Distance to rivers, major, minor creeks.</td>
<td>Euclidean distance to rivers, major, minor creeks. 25 m raster data.</td>
</tr>
<tr>
<td></td>
<td>(Fiji Land Information System). Original 1:50K topographic maps [23].</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Roads</td>
<td>Fiji Ministry of Lands and Resources. Digital data from FLIS (Fiji</td>
<td>Road density.</td>
<td>Number of roads per sq km within 1 km radius. 25 m raster data.</td>
</tr>
<tr>
<td></td>
<td>Land Information System). Original 1:50K topographic maps.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Land use/</td>
<td>Ministry of Agriculture. Digital data from Secretariat of the Pacific</td>
<td>Multiple land use/cover types.</td>
<td>Visual interpretation of satellite imagery. 25 m raster data.</td>
</tr>
<tr>
<td>cover</td>
<td>(SPC). Circa 2010.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Climate</td>
<td>Landcare Research Institute [25].</td>
<td>Annual, maximum, minimum, and average temperature and rainfall.</td>
<td>Spatially interpolated climate data meteorological station data from 1971–2000. 100 m</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>raster data.</td>
</tr>
<tr>
<td>Elevation</td>
<td>Landcare Research Institute [3].</td>
<td>Altitude and slope</td>
<td>Elevation derived from 20 m contours. 25 m raster data.</td>
</tr>
<tr>
<td>Census</td>
<td>Fiji National Census 2007 [20].</td>
<td>Multiple measures of education attainment, house construction, employment, ethnicity,</td>
<td>Census variables available at the Enumeration Area level (~80–120 households). Vector</td>
</tr>
<tr>
<td></td>
<td></td>
<td>and other sociodemographic factors.</td>
<td>count data converted to proportions.</td>
</tr>
<tr>
<td>Economic</td>
<td>World Bank (2011) report [26].</td>
<td>Poverty rate (% of population below poverty line) and poverty gap (how far on</td>
<td>Poverty rates estimated using small area estimation method. Vector data at the Tikina</td>
</tr>
<tr>
<td>status</td>
<td></td>
<td>average people are from the poverty line)</td>
<td>level.</td>
</tr>
<tr>
<td>Livestock</td>
<td>Fiji Ministry of Agriculture. Fiji National Agricultural Census 2009</td>
<td>Number of farms and farm animals by species: cattle, commercial beef, dairy,</td>
<td>Density per sq km calculated by Tikina (not available for all Tikinas).</td>
</tr>
<tr>
<td></td>
<td>[27].</td>
<td>subsistence beef, etc.</td>
<td></td>
</tr>
</tbody>
</table>

Census [20], and data on poverty rates from the 2011 World Bank Report [26]. Livestock data were provided by the Fiji Ministry of Agriculture’s 2009 National Agricultural Census [27]. All geospatial data were georeferenced to the Fiji Map Grid 1986 coordinate system. Table 1 provides a summary of the environmental, census, socio-demographic and livestock data used in the study. The five geographic regions used in this study and examples of the geo-referenced data are shown in Fig 3.
Fig 3. Fiji geography, and examples of environmental and census data used: a) Divisions and ‘regions’ included in the study, major rivers; b) altitude; c) rainfall; d) total cattle density; e) poverty rate; f) proportion of households with metered (treated) water at home. See Table 1 for data sources.

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and community-level data include risk factors that are common to all inhabitants of a household and community respectively.

- **Individual-level data.** Potential risk factors for leptospirosis were assessed using questionnaire-based interviews, including demographics, occupation, recreational activities, contact with animals, education, and knowledge about leptospirosis.

- **Household-level data.** Information on household income, house construction, access to utilities (toilets, water, sewage, electricity), and the presence of animals and crops around the home were obtained through questionnaires. In addition, data on environmental attributes (including rainfall, temperature, elevation, land cover, soil type, and distance to rivers) at household locations were extracted or calculated using geographic information systems (GIS) as described above.

- **Community-level data.** Community type, urban/rural settings, and the presence of animal species in each community were ascertained through questionnaires. Census and agricultural data were extracted or calculated using the geospatial databases described in Table 1. Data were available at the enumeration area resolution (~80–120 households) for a variety of socioeconomic and demographic measures, including the proportions of households with metered water, toilets, electricity, and sewage services; population ethnicity, level of educational attainment, and reliance on subsistence farming as the main source of income. At the Tikina level, data were available on World Bank estimates of poverty measures, and census of animal populations conducted by the Fiji Ministry of Agriculture.

**Maps**

Maps were produced to show the locations of communities that participated in the field study, and the observed community-level seroprevalence in 2013. Although all household GPS locations were recorded, only community-level seroprevalence were depicted on maps to protect the identity of participants. Locations of communities were mapped to their median centre, calculated as the location nearest to all sampled households in the community while minimizing the effects of outliers.

**Serological analysis**

Blood samples were processed in Fiji, and frozen sera transported to Australia for serological analysis. Microscopic agglutination tests (MAT) were used to detect anti-*Leptospira* antibodies, and determine the putative serogroups associated with infections. The MAT is the reference serological test recommended by the WHO and the International Committee on Systematic Bacteriology (Subcommittee on the Taxonomy of *Leptospira*) [28,29]. Serological analyses were conducted at the WHO/FAO/OIE Collaborating Centre for Reference & Research on Leptospirosis in Brisbane, Australia. Based on the laboratory’s knowledge of the epidemiology of leptospiral serovars in the South Pacific, 21 pathogenic serovars (see S1 Appendix) were selected for the initial MAT panel for this study, and samples were tested at dilutions from 1:50 to 1:3200. The 21-serovar panel was used to test a random selection of ~10% of total samples to determine the most common serogroups responsible for infections. In addition, the 21-serovar panel was used to test 199 *Leptospira* ELISA-positive samples collected from patients with suspected clinical leptospirosis in Fiji in 2012 and 2013 to ensure that the most common serogroups associated with clinical infections were included in the final panel. Based on the MAT results from the two sets of sera, six serovars were chosen for the final panel used to test the remaining samples from this study (S1 Appendix): *Leptospira interrogans* serovars Pohnpei...
(serogroup Australis), Australis (serogroup Australis), Canicola (serogroup Canicola), Copen-
hageni (serogroup Icterohaemorrhagiae), Hardjo (serogroup Sejroe), and \textit{Leptospira borgpeter-
senii} serovar Ballum (serogroup Ballum).

The MAT assay is expensive and time-consuming, and the described strategy to limit the
number of serovars included in the final panel resulted in reduced project costs. Considering
that one dominant serovar was identified in the preliminary tests, it was determined that the
smaller MAT panel was unlikely to have significant impact on the overall epidemiological find-
ings. MAT titres of $\geq 1:50$ were considered reactive or seropositive, and indicative of recent or
past infection. For samples that reacted to multiple serovars within a serogroup, the serovar
associated with the highest titre was considered to be the reacting serovar. Samples that reacted
to serovars in more than one serogroup were recorded as reacting to multiple serovars.
Although serogroups are no longer used in the taxonomic classification of serovars, they
remain useful for laboratory purposes and epidemiological comparisons.

Statistical analysis

An overview of the statistical analyses is shown in \textbf{Fig 2}. The outcome measure used was sero-
positive reactions to any of the six serovars included in the final MAT panel. Firstly, crude asso-
ciations between the independent variables and the outcome measure were obtained by
univariable logistic regression. Independent variables associated with the outcome by a likeli-
hood ratio test (LRT) $p$-value of $<0.2$ were then subjected to a stepwise backward elimination
process ($p<0.05$) to select the final set of independent variables for the multivariable logistic
regression models. In addition, the possible presence of effect modification in the multivariable
modelling was investigated using the LRT. This was assessed using interaction terms, which
consisted of all independent variables found to be significant in the univariable analysis. Inter-
action terms were added separately to the analyses to determine their joint effect on the out-
come measure. Multilevel hierarchical modelling was used to take into account the clustering
of participants, and allowed for correlation of observations by \textit{region} ($n=5$), \textit{community}
($n=81$), and \textit{household} (up to 3 participants per household in 8a) as random effects. Intra-
cluster correlation coefficients (ICCs) with corresponding 95% confidence intervals were
obtained from final multivariable models. Biological plausibility and collinearity between vari-
ables were taken into account when selecting the variables to be retained in the final models.
For example, if we observed strong collinearity between poverty rate and % of households in
the community with electricity supply, poverty rate would be chosen for the final model
because of more direct relationship to exposure risks.

Two multivariable models were built:

- **Model A.** Used independent variables where data could be ascertained by questioning an indi-
  vidual, and included primarily individual-level variables, but also some household-level and
  community-level variables. Model A was designed to assess the risk of infection for an individ-
  ual, e.g. for producing predictive risk charts to graphically depict the combined effects of vari-
  ables in determining overall seroprevalence. The charts are designed for use in clinical settings,
  and are similar to cardiovascular risk charts used to predict the risk of cardiac events based on
  combinations of risk factors such as blood pressure, diabetes, smoking, and cholesterol levels.

- **Model B.** Used independent variables derived from geospatial and other national databases
described in \textbf{Table 1}, without using the field data collected in this study. Model B included
  household-level and community-level variables only, and was designed to assess the risk of
  infection at geographic locations, e.g. estimating community-level seroprevalence, identifying
  hotspots and producing predictive risk maps.
Independent variables found to be statistically significant on multivariable regression analyses were reported. Adjusted odds ratios (OR) with 95% confidence intervals obtained from regression coefficients were used to quantify associations between the independent and outcome variables. In addition, univariate results of variables associated with animal exposure and contact were reported. Statistical significance was considered at \( p < 0.05 \) and two-sided. Data analysis was performed using STATA 13 (StataCorp, 2013). Model fit was assessed using the Hosmer-Lemeshow test [30], while relative predictive performance was undertaken using the area under the receiver operating curve (AUC) was calculated for each model and compared for statistical differences. An AUC of 0.7 was deemed to indicate an adequate predictive ability of the model. Akaike information criterion (AIC) and Bayesian information criterion (BIC) were reported for the final models.

**Results**

**Study population**

A total of 2152 participants from 1922 households in 81 communities on the three main islands of Fiji were included in the study. The age of participants ranged from 1 to 90 years (mean 33.6, SD 19.8), and 985 (45.8%) were males. The age and sex distribution of participants are shown in Fig 4. The study included 662 participants from the Central Division (28 communities), 453 from Ba (10 communities), 520 from other parts of the Western Division (21 communities), 261 from Taveuni (11 communities), and 256 from Vanua Levu (11 communities).
Table 2. *Leptospira* seroprevalence by age, gender, ethnicity, community types, and region.

<table>
<thead>
<tr>
<th>Variables</th>
<th>No of participants</th>
<th>Reactive MATs*</th>
<th>Sero-prevalence (%)</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total sampled</td>
<td>2152</td>
<td>417</td>
<td>19.4%</td>
<td>17.7–21.1%</td>
</tr>
<tr>
<td><strong>Gender</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>985</td>
<td>234</td>
<td>23.8%</td>
<td>21.1–26.5%</td>
</tr>
<tr>
<td>Female</td>
<td>1160</td>
<td>182</td>
<td>15.7%</td>
<td>13.6–17.9%</td>
</tr>
<tr>
<td><strong>Age groups (years)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–9</td>
<td>256</td>
<td>21</td>
<td>8.2%</td>
<td>5.1–12.3%</td>
</tr>
<tr>
<td>10–19</td>
<td>362</td>
<td>69</td>
<td>19.1%</td>
<td>15.1–23.5%</td>
</tr>
<tr>
<td>20–29</td>
<td>387</td>
<td>101</td>
<td>26.1%</td>
<td>21.8–30.8%</td>
</tr>
<tr>
<td>30–39</td>
<td>340</td>
<td>61</td>
<td>17.9%</td>
<td>14.0–22.4%</td>
</tr>
<tr>
<td>40–49</td>
<td>279</td>
<td>63</td>
<td>22.6%</td>
<td>17.8–27.9%</td>
</tr>
<tr>
<td>50–59</td>
<td>263</td>
<td>50</td>
<td>19.0%</td>
<td>14.5–24.3%</td>
</tr>
<tr>
<td>≥ 60</td>
<td>263</td>
<td>52</td>
<td>19.8%</td>
<td>15.1–25.1%</td>
</tr>
<tr>
<td><strong>Ethnicity</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indo-Fijian</td>
<td>459</td>
<td>34</td>
<td>7.4%</td>
<td>5.2–10.2%</td>
</tr>
<tr>
<td>iTaukei</td>
<td>1651</td>
<td>374</td>
<td>22.7%</td>
<td>20.7–24.7%</td>
</tr>
<tr>
<td>Other</td>
<td>39</td>
<td>8</td>
<td>20.5%</td>
<td>9.3–36.5%</td>
</tr>
<tr>
<td><strong>Community type</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Private residential</td>
<td>502</td>
<td>44</td>
<td>8.8%</td>
<td>6.4–11.6%</td>
</tr>
<tr>
<td>Settlement (Indo-Fijian)</td>
<td>103</td>
<td>18</td>
<td>17.5%</td>
<td>10.7–26.2%</td>
</tr>
<tr>
<td>Settlement (mixed ethnicity)</td>
<td>511</td>
<td>91</td>
<td>17.8%</td>
<td>14.6–21.4%</td>
</tr>
<tr>
<td>Village</td>
<td>1036</td>
<td>264</td>
<td>25.5%</td>
<td>22.9–28.3%</td>
</tr>
<tr>
<td><strong>Urban/Rural</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban</td>
<td>579</td>
<td>64</td>
<td>11.1%</td>
<td>8.6–13.9%</td>
</tr>
<tr>
<td>Peri-urban</td>
<td>287</td>
<td>44</td>
<td>15.3%</td>
<td>11.4–20.0%</td>
</tr>
<tr>
<td>Rural</td>
<td>1286</td>
<td>309</td>
<td>24.0%</td>
<td>21.7–26.5%</td>
</tr>
<tr>
<td><strong>Region</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central Division</td>
<td>662</td>
<td>107</td>
<td>16.2%</td>
<td>13.4–19.2%</td>
</tr>
<tr>
<td>Western Division–Ba</td>
<td>453</td>
<td>82</td>
<td>18.1%</td>
<td>14.7–22.0%</td>
</tr>
<tr>
<td>Western Division–Other</td>
<td>520</td>
<td>94</td>
<td>18.1%</td>
<td>14.9–21.7%</td>
</tr>
<tr>
<td>Northern Division–Taveuni</td>
<td>261</td>
<td>59</td>
<td>22.6%</td>
<td>17.7–28.2%</td>
</tr>
<tr>
<td>Northern Division–Vanua Levu</td>
<td>256</td>
<td>75</td>
<td>29.3%</td>
<td>23.8–35.3%</td>
</tr>
</tbody>
</table>

*Reactive MAT defined at titre of ≥1:50 for one or more serovars used in the 6-serovar panel

| doi:10.1371/journal.pntd.0004405.t002 |

(Table 2). One of the selected communities in Taveuni was not included because of logistic constraints.

**Seroprevalence and serovars**

Details of the 21 serovars included in the screening panel and the 6 serovars included in the final panel are shown in S1 Appendix, together with the seroprevalence of the initial 198 randomly selected samples from this study and the 199 *Leptospira* ELISA-positive samples from patients with suspected leptospirosis from April 2012 to November 2013. The six serovars included in the final MAT panel accounted for 86.7% of reactive tests: *Leptospira interrogans* serovars Pohnpei, Australis, Canicola, Copenhageni, and Hardjo; and *Leptospira borgpetersenii* serovar Ballum. Using the 6-serovar panel, the overall seroprevalence was 19.4% (95% CI 17.7%–21.1%), with 417 participants having reactive MATs to at least one serovar. One
predominant serovar, Pohnpei, accounted for 351 (84.2%; 95% CI 80.3%–87.5%) of reactive MATs. A total of 63 participants had MAT titres of ≥1:400 (47 for serovar Pohnpei, and 16 for other serovars), the cutoff used by our laboratory to indicate an acute infection. The distribution of MAT titres for Pohnpei and other serovars is shown in Fig 5.

Table 2 shows that there were significant differences in seroprevalence by age, gender, ethnicity, community types, and region. Community-level seroprevalence ranged from 0% to 60%, and are shown on the maps in Fig 6a to 6d. Variations in seropositive reactions to each serovar by age groups and region of residence are shown in Fig 7a & 7b respectively.

Risk factor analysis and multivariable models

A total of 118 independent variables were assessed on univariate analysis: 75 variables obtained from questionnaires, and 43 derived using GIS from the sources described in Table 1. Independent variables included 31 individual-level, 38 household-level, and 49 community-level risk factors described above. S2 Appendix provides a list of the independent variables assessed at the univariate level. Variables statistically significant on univariate analyses were considered for the multivariable models, and included 19 individual-level, 21 household-level, and 25
community-level risk factors. Due to statistical significance not be reached, no interaction effect was included in the multivariable modelling.

Multivariable Model A (using variables where data could be ascertained by questioning an individual) included five variables that were independently associated with the presence of *Leptospira* antibodies, with an AUC of 0.7 (Table 3) including: male gender (OR 1.55 compared to females), iTaukei ethnicity (OR 3.51 compared to Indo-Fijians), living in settlements and villages (OR 2.13 and 1.64 respectively compared to urban residential areas), not having metered water at home (OR 1.52), and working outdoors (OR 1.64 compared to working indoors). Of the 434 participants who worked outdoors, 378 (87%) were full- or part-time farmers, indicating that outdoor work in Fiji is predominantly related to farming.

Multivariable Model B (using only variables derived from geospatial and other national databases) included six variables that were independently associated with the presence of *Leptospira* antibodies, with an AUC of 0.7 (Table 4) including: living in rural areas (OR 1.43...
Fig 7. Percentage of positive MAT reactions associated with each of the 6 serovars included in the final panel by: a) age groups, and b) regions. Positive MAT reactions defined as titre of ≥ 1:50.

doi:10.1371/journal.pntd.0004405.g007
compared to living in urban or peri-urban areas), poverty rate ≥ 40% (OR 1.74), living <100m from a river or major creek (OR 1.41), presence of pigs in the community (OR 1.54), total cattle density in the Tikina (OR 1.04 per head of cattle per square km), and high maximum rainfall in the wettest month (OR 1.003 per mm of rain). Total cattle density (includes both commercial and subsistence livestock) ranged from 0.11 to 31.48 head of cattle per square km (mean 8.96, SD 5.31), and maximum rainfall in the wettest month ranged from 275 to 789mm (mean 375.02, SD 56.94). A similar multivariable model using total dairy farm density instead of total cattle density performed better than the final model, but data on dairy farm density were only available for 57 of the 81 (70.4%) communities included in our study, and this variable was therefore not selected for the final Model B.

Collection of biological samples from animals was outside of the scope of this study, but study questionnaires included detailed information about contact with animals (rodents, mongoose, pets, and livestock) at home and the presence of animals in the community. A number of animal-related exposures were significantly associated with the presence of *Leptospira* antibodies on univariable analysis (Table 5). The presence of rats, mice, and mongoose around the home was not significantly associated with seroprevalence, but higher infection rates were found in participants who reported physical contact with rats or mice (OR 1.58) and mongoose (OR 1.81). Table 5 shows that many Fijians have livestock animals at home and in the community. The presence of each livestock species was associated with a higher infection rates on

| Table 3. Variables significantly associated with positive MAT for *Leptospira* on univariable and multivariable analysis—Model A* (individual-level variables). |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Variables       | No of subjects  | Reactive MAT*   | Sero-prevalence (%) | Univariable Odds Ratio (95% CI) | Adjusted Odds Ratio (95% CI) | p value* |
| Total sampled   | 2152            | 417             | 19.4%             | 1                            | 1                             | 0.003     |
| Gender          |                 |                 |                  |                               |                               |           |
| Female          | 1160            | 182             | 15.7%            | 1.67 (1.35–2.08)              | 1.55 (1.16–2.08)              |           |
| Male            | 985             | 234             | 23.8%            |                               |                               |           |
| Ethnic group    |                 |                 |                  |                               |                               |           |
| Indo-Fijian     | 459             | 34              | 7.4%             | 1                            | 1                             | <0.001    |
| iTaukei         | 1651            | 374             | 22.7%            | 3.66 (2.53–5.29)              | 3.51 (2.23–5.54)              |           |
| Other           | 39              | 8               | 20.5%            | 3.23 (1.38–7.56)              | 2.32 (0.82–6.58)              | 0.114     |
| Community type  |                 |                 |                  |                               |                               |           |
| Urban residential | 502          | 44              | 8.8%             | 1                            | 1                             | <0.001    |
| Settlement      | 614             | 109             | 17.8%            | 2.25 (1.55–3.26)              | 2.13 (1.41–3.21)              |           |
| Village         | 1036            | 264             | 25.5%            | 3.56 (2.54–5.00)              | 1.64 (1.08–2.51)              | 0.021     |
| Metered water available at home | | | | | | |
| Yes             | 1412            | 221             | 15.7%            | 1.92 (1.54–2.39)              | 1.52 (1.14–2.03)              | 0.004     |
| No              | 720             | 189             | 26.3%            |                               |                               |           |
| Work location   |                 |                 |                  |                               |                               |           |
| Indoors         | 832             | 106             | 12.7%            | 1.63 (1.23–2.17)              | 1.65 (1.23–2.20)              | 0.001     |
| Mixed indoors/ outdoors | 639 | 123 | 19.2% | 1.63 (1.23–2.17) | 1.65 (1.23–2.20) | 0.001 |
| Outdoors        | 434             | 119             | 27.4%            | 2.59 (1.93–3.47)              | 1.64 (1.15–2.34)              | 0.006     |

* Model goodness of fit: AIC 1675.9, BIC 1731.3, df 10.
* Reactive MAT defined at titre of ≥1:50 for one or more serovars used in the 6-serovar panel
* p value for adjusted odds ratios, multivariable model

doi:10.1371/journal.pntd.0004405.t003
univariable analysis, but only the presence of pigs in the community was significant on multi-
variable analysis, and included in Model B.

Multilevel hierarchical models were built to take into account spatial correlation of data: i) defining region and community as random effects, using the entire dataset, and ii) defining community and household as random effects, using Ba data only. The results of multilevel models were not statistically different to the results of multivariable Model A using all data (chi2 = 0.01, p = 0.99), Model A using Ba data (chi2 = 0.00, p = 1.00), Model B using all data (chi2 = 0.01, p = 0.99), or Model B using Ba data (chi2 = 0.00, p = 1.00). For all multilevel models, intra-class correlation coefficients were <0.01 and odds ratio estimates were very similar to the reported models. Results for the multilevel models were therefore not reported here.

Seroprevalence estimation chart using Model A

Fig 8 shows a seroprevalence estimation chart that incorporates individual-level variables to show the combined effects of multiple independent risk factors on the prevalence of infection. Estimated seroprevalence were based on the five variables used in Model A. For example, the chart shows a range of seroprevalence from 2.0% for female Indo-Fijians who live in urban residential areas, have metered water at home, and work indoors; to 39.4% for male iTaukei who

Table 4. Variables significantly associated with positive MAT for *Leptospira* on univariable and multivariable analyses–Model B^ (environmental and census variables).

<table>
<thead>
<tr>
<th>Variables</th>
<th>No of subjects</th>
<th>Reactive MAT*</th>
<th>Sero-prevalence (%)</th>
<th>Univariable Odds Ratio (95% CI)</th>
<th>Adjusted Odds Ratio (95% CI)</th>
<th>p value^#</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urban/Rural</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban/ Peri-urban</td>
<td>866</td>
<td>108</td>
<td>12.5%</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Rural</td>
<td>1286</td>
<td>309</td>
<td>24.0%</td>
<td>2.22 (1.75–2.82)</td>
<td>1.43 (1.07–1.91)</td>
<td>0.016</td>
</tr>
<tr>
<td>Poverty rate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 40%</td>
<td>1277</td>
<td>187</td>
<td>14.6%</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>≥ 40%</td>
<td>875</td>
<td>230</td>
<td>26.3%</td>
<td>2.08 (1.67–2.58)</td>
<td>1.74 (1.31–2.31)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Distance between home and river or major creek</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt; 100m</td>
<td>1590</td>
<td>279</td>
<td>17.6%</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>≤ 100m</td>
<td>456</td>
<td>115</td>
<td>25.2%</td>
<td>1.58 (1.24–2.03)</td>
<td>1.41 (1.09–1.83)</td>
<td>0.009</td>
</tr>
<tr>
<td>Presence of pigs in community</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>1587</td>
<td>266</td>
<td>16.8%</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>561</td>
<td>150</td>
<td>26.7%</td>
<td>1.81 (1.44–2.28)</td>
<td>1.54 (1.21–1.98)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Mean (Standard deviation)

| Total cattle density in Tikina^§ (per head of cattle per sq km) | Seronegative subjects | 8.86 (5.18) | Seropositive subjects | 9.38 (5.83) |
| Maximum rainfall in wettest month (per mm)                      | Seronegative subjects | 372.83 (50.05) | Seropositive subjects | 384.18 (79.01) |

^Model goodness of fit: AIC 1924.3, BIC 1963.6, df 7.
*Reactive MAT defined at titre of ≥1:50 for one or more serovars used in the 6-serovar panel
# p value for adjusted odds ratios, multivariable model
§ Includes both commercial and subsistence cattle

doi:10.1371/journal.pntd.0004405.t004
live in mixed ethnic settlements, do not have metered water and home, and work outdoors. It is uncommon for Indo-Fijians to live in villages or for iTaukei to live in Indo-Fijian settlements, and results were therefore not shown for these scenarios.

**Discussion**

Our study identified a high risk of human leptospirosis infection in Fiji, with an overall seroprevalence of 19.4% using a 6-serovar MAT panel. One dominant serovar, Pohnpei, was associated with 84.2% of reactive MATs. The serovar was originally isolated from rodents and pigs during an animal leptospirosis study in the island of Pohnpei in the Federated States of Micronesia [31], and has been found to be an important cause of human infections [32]. Seroprevalence varied significantly between the five regions in our study, and ranged from 16.2% in the Central Division to 29.3% in Vanua Levu in the Northern Division. Community-level seroprevalence also varied significantly from 0% to 60% in the 81 communities included in our study. These findings indicate marked geographic variation in infection risk in Fiji and the presence of hotspots where disease transmission is more intense.

Globally, reported leptospirosis seroprevalence vary significantly between and within countries, based on environmental settings, behavioural risk factors, and socio-demographics; our results corroborate these findings. To put the Fiji results into a global context, examples of seroprevalence reported from known high risk settings [2,4] such as urban slums, tropical islands, and flood risk areas include 15.4% in an urban slum in Brazil [33], 37% of healthy adult males in the Seychelles [34], 18.8% in the Mekong delta in Vietnam [35], and 23.9% and 38.2% in flood-prone areas in Laos and Bangladesh respectively [36,37]. As found in Fiji, small-scale variations in seroprevalence within countries and differences between occupational groups have been reported. In American Samoa (a group of remote islands in the south Pacific), a community-based study reported an overall seroprevalence of 15.5% [8] and significant

**Table 5. Associations between positive MAT for Leptospira and animal exposure at home and in the community.**

<table>
<thead>
<tr>
<th>Questions related to animal exposure and contact</th>
<th>Number of subjects who answered ‘yes’</th>
<th>% of subjects who answered ‘yes’</th>
<th>Univariable Odds Ratio (95% CI)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seen rats or mice at or around your home</td>
<td>1844</td>
<td>85.9%</td>
<td>1.16 (0.84–1.59)</td>
<td>0.371</td>
</tr>
<tr>
<td>Been in physical contact with rats or mice</td>
<td>323</td>
<td>15.3%</td>
<td>1.58 (1.20–2.09)</td>
<td>0.001</td>
</tr>
<tr>
<td>Seen mongooses at or around your home</td>
<td>1655</td>
<td>77.1%</td>
<td>1.08 (0.83–1.39)</td>
<td>0.574</td>
</tr>
<tr>
<td>Been in physical contact with mongooses</td>
<td>135</td>
<td>6.5%</td>
<td>1.81 (1.23–2.68)</td>
<td>0.003</td>
</tr>
<tr>
<td>Pigs at your home or in your garden</td>
<td>230</td>
<td>10.7%</td>
<td>1.55 (1.23–2.12)</td>
<td>0.007</td>
</tr>
<tr>
<td>Pigs in your community</td>
<td>561</td>
<td>26.1%</td>
<td>1.81 (1.44–2.28)</td>
<td>0.000</td>
</tr>
<tr>
<td>Cows at your home or in your garden</td>
<td>284</td>
<td>13.2%</td>
<td>1.53 (1.15–2.05)</td>
<td>0.004</td>
</tr>
<tr>
<td>Cows in your community</td>
<td>481</td>
<td>22.4%</td>
<td>1.52 (1.19–1.93)</td>
<td>0.001</td>
</tr>
<tr>
<td>Horses at your home or in your garden</td>
<td>200</td>
<td>9.3%</td>
<td>1.53 (1.09–2.14)</td>
<td>0.013</td>
</tr>
<tr>
<td>Horses in your community</td>
<td>377</td>
<td>17.6%</td>
<td>1.55 (1.19–2.01)</td>
<td>0.001</td>
</tr>
<tr>
<td>Are there goats at your home or in your garden?</td>
<td>107</td>
<td>5.0%</td>
<td>1.08 (0.67–1.75)</td>
<td>0.749</td>
</tr>
<tr>
<td>Goats in your community</td>
<td>242</td>
<td>11.3%</td>
<td>1.47 (1.08–2.01)</td>
<td>0.015</td>
</tr>
<tr>
<td>Chickens at your home or in your garden</td>
<td>431</td>
<td>20.1%</td>
<td>1.21 (0.93–1.57)</td>
<td>0.152</td>
</tr>
<tr>
<td>Chickens in your community</td>
<td>819</td>
<td>38.1%</td>
<td>1.39 (1.12–1.72)</td>
<td>0.003</td>
</tr>
<tr>
<td>Dogs at your home or in your garden</td>
<td>645</td>
<td>30.0%</td>
<td>1.00 (0.79–1.26)</td>
<td>0.992</td>
</tr>
<tr>
<td>Dogs in your community</td>
<td>998</td>
<td>46.5%</td>
<td>1.25 (1.01–1.55)</td>
<td>0.041</td>
</tr>
<tr>
<td>Cats at your home or in your garden</td>
<td>355</td>
<td>16.5%</td>
<td>0.78 (0.58–1.06)</td>
<td>0.115</td>
</tr>
<tr>
<td>Cats in your community</td>
<td>830</td>
<td>38.6%</td>
<td>1.38 (1.11–1.72)</td>
<td>0.003</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pntd.0004405.t005
variation between islands with different environments, and between areas with different population density [38]. In Peru, seroprevalence varied from 28.0% in the Amazonian city of Iquitos and 16.5% in the surrounding villages (wet tropics), to 0.7% in a desert shantytown near Lima [39]. In the Andaman Islands, a study of high-risk populations found seroprevalence of 62.5% in agricultural workers, 39.4% in sewage workers, 37.5% in animal handlers, and 30.0% in butchers [40]. In contrast, a study of healthy blood donors in an area of high leptospirosis incidence in northern Queensland in Australia found a seroprevalence of only 1.4% [41,42].

Our study found that individual-level factors were important predictors of leptospirosis infection risk in Fiji. Model A shows that gender, ethnicity, community type, availability of water at home, and work location were independently associated with the presence of *Leptospira* antibodies. Higher infection rates in males corroborates findings in the majority of leptospirosis studies around the world, and is likely to be associated with higher frequency of outdoor activities as well as higher risk occupational and recreational exposures. Reasons for the marked difference in seroprevalence between the two main ethnic groups in Fiji are unclear, but could be related to differences in genetic susceptibility or behaviours that were not elucidated by our questionnaire, e.g. differences in animal husbandry or slaughtering practices related to religion or culture. Further studies are required to explain the disparate risk between ethnic groups. Seroprevalence in villages was significantly higher than in urban residential communities.
areas or settlements, and is likely the result of more intimate contact with the natural environment and domestic animals. In our study, working outdoors was associated with a higher risk of infection, and the majority of outdoor work in Fiji involves farming. Agriculture is an important part of Fiji’s economy, and apart from the livestock industry, there is commercial farming of a range of crops include sugarcane, coconut, copra, and a wide variety of fruits and vegetables. Occupational exposure in the agricultural industry is therefore likely to be an important source of leptospirosis infection in Fiji.

Of note, three of the predictors included in our final multivariable models were related to water: the availability of metered (treated) water at home (Model A), distance between the home and the closest river or major creek (Model B), and maximum rainfall in the wettest month (Model B). Considering that Leptospira can survive for weeks to months in fresh water, and are efficiently carried and disseminated by water (e.g. flooding, flowing downstream in rivers), the findings were not unexpected. Lack of metered water at home and proximity to rivers are likely to be associated with higher levels of contact with untreated freshwater, e.g. using rivers for bathing, cleaning, swimming, and recreational activities. Furthermore, poor access to water at home is generally associated with poverty (discussed below), and also influences personal hygiene, e.g. the ability to clean and wash after working outdoors, or after contact with mud, contaminated water, or animals. Two of the water-related predictors (distance to river or major creek and maximum rainfall in the wettest month) are also proxy measures of flooding risk. As seen with the post-flood leptospirosis outbreaks in 2012, flooding is an important driver of transmission in Fiji, as it is in many parts of the world.

Two of the predictors in Model B relate to livestock exposure: total cattle density in the Tikina and presence of pigs in the community. Data on cattle density in Tikinas includes both commercial and subsistence farming, and varied from < 1 to over 30 heads of cattle per square km. Infection risk could be related to direct occupational contact with cattle, or through more general contamination of the environment (especially rivers) with cattle urine. As shown in Table 5, many households in Fiji keep subsistence livestock. Backyard piggeries are commonly found in communities in Fiji and other Pacific Islands, and are usually small pens with less than 10 pigs. The pens are often built on the edge of rivers and streams to allow convenient drainage of waste, but unfortunately also lead to contamination of freshwater at that community as well as further downstream. In American Samoa, similar backyard piggeries have been associated with the risk of human leptospirosis infection [8,43]. Dairy farmers are known to be at high risk for leptospirosis in many parts of the world because of close contact with cattle, and exposure to urine during milking. In our study, high density of dairy farms was strongly associated with infection risk, but was not included as a variable in the final model because data were only available for ~70% of the Tikinas in our study. As more data on dairy farms become available, associations with leptospirosis risk could be further explored and model performance potentially improved. Commercial dairy and beef farming could potentially intensify in the future with population growth, and increase the risk of leptospirosis.

Model B also shows that leptospirosis is a disease of poverty in Fiji and disproportionately affects the poorest. Leptospirosis has been associated with poverty in diverse settings around the world, including Brazilian and Indian urban slums [33,44,45], Peruvian Amazon [39], and areas of poor socioeconomic status in the USA and Europe [46,47]. Furthermore, the combination of poverty, livestock keeping, and global climate change are important drivers of zoonotic diseases transmission [48]. In our study, participants living in communities with high poverty rates (defined as ≥40% of households in the community) had almost twice the infection rate compared to other communities, independent of the other predictors in Model B. As discussed above, poor access to metered (treated) water at home was associated with a higher risk of infection for many reasons, and is also a proxy measure of socioeconomic status.
Although serovar Pohnpei was associated for 84.2% of reactive MATs, there were differences in serovar distribution by age and by region of residence, suggesting that the relative importance of animal species in disease transmission varies between subgroups. Variation in risk factors between age groups likely relates to age-specific behaviours, e.g. young children spend more time playing around the home, and have closer contact with pets and soil; teenagers have more frequent recreational freshwater contact from swimming in rivers and waterfalls; and adults have more intense contact with livestock through occupational exposure and managing animals at home. Variation in risk factors between regions likely relates to differences in environmental settings, with proportionately greater urbanization in the Central division, and more farming in the other regions. For example, rodents could be more important in transmission cycles in urban and peri-urban areas, and livestock more important in rural areas.

Many of the risk factors and environmental drivers identified in our study provide significant cause for concern about future risk of leptospirosis in Fiji, as well as other Pacific Islands with similar environments. Population growth is typically associated with agricultural intensification, leading to increase in livestock numbers (both commercial and subsistence) and occupational exposure. With global climate change, extreme weather events and flooding are predicted to become more frequent and intense in the Pacific Islands. Rapid population growth in developing countries is often associated with urban and peri-urban slums where diseases of poverty proliferate. Although our study found that leptospirosis seroprevalence was lower in urban areas, poverty rate was a significant risk factor independent of urban or rural settings. Climate change, flooding, population growth, urbanization, and agricultural intensification may independently, or potentially synergistically, lead to enhanced leptospirosis transmission in Fiji [3].

The findings should be considered in light of the study’s limitations. Limitations of the MAT have been well documented; the test is considered to be serogroup rather than serovar specific, cross-reactions occur between serovars within a serogroup, and complex paradoxical reactions could occur in persons who have had previous infections [29]. Despite these limitations, the MAT is considered the gold standard test for identifying putative serogroups and serovars when isolates are not available [28]. Isolates of leptospires would be required to definitively confirm the serovars circulating in Fiji. Due to budgetary reasons, our study used a 6- rather than 21-serovar MAT panel to test the majority of samples. If the larger panel was used, additional less-common serovars may have been detected. However, the 6 serovars selected included the most reactive serovars when 198 randomly selected samples from this study were tested against the full 21-serovar panel; the 6 serovars selected accounted for 86.7% of the reactive samples, and one dominant serovar (Pohnpei) accounted for 65.9% of reactive samples. The reduced MAT panel size could have underestimated the overall seroprevalence by a factor of 0.13 compared to a 21-serovar panel, but unlikely to have significantly influenced the overall epidemiological patterns reported here because one serovar dominated the reactive MATs, and our data analyses in this paper were not stratified by serovars.

Our study measured antibodies to *Leptospira* to identify evidence of prior infection. However, many leptospirosis infections do not result in any apparent illness and are of no clinical significance. The severity of clinical disease depends on many factors including pathogen virulence and the individual’s immune status, comorbidities, and age [49]. Serovar Pohnpei, the serovar associated with 84.2% of MAT-reactive cases, has been reported as an important cause of overt clinical disease in the Federated States of Micronesia [32], suggesting the findings in this study are applicable not only to infection risk but also clinical illness. However, there are currently no available data on the proportion of serovar Pohnpei infections that result in clinical disease or severe complications.

Future studies could further improve our understanding of leptospirosis transmission in Fiji by examining serovar-specific risk factors; identifying the most important exposures in
different subgroups such as age groups, gender, ethnic groups, and community types; deter-
miming the relative importance of livestock, rodents, pets and wildlife in transmitting leptospiri-
rosis to humans; and developing models to determine transmission (causal) pathways rather
than just epidemiological links. For environmental and census variables, we used data at the
place of residence, but infections could also have occurred at work or elsewhere. Future studies
that focus specifically on work-related activities would provide more insight into the impor-
tance of occupational exposures in Fiji. The performances of models were partly determined by
the accuracy of available environmental, census, and livestock data, and models could be
updated and improved as more data become available. Models based on environmental factors,
such as Model B, could be used to produce predictive risk maps for the whole of Fiji.

In summary, our study found that risk factors and drivers for human leptospirosis infection
in Fiji are complex and multifactorial, and include climate, the natural environment, livestock
(both subsistence and commercial), living conditions, socioeconomic status, demographics and
individual behaviour. Some of these factors corroborate findings previously reported in differ-
ent settings (e.g. male gender, working outdoors), but other factors appear to be specific to the
cultural and environmental settings in Fiji, including ethnicity and presence of pigs in commu-
nities. By using an integrated eco-epidemiological approach and including a wide range of data
sources in our analyses, we were able to quantify the relative importance of risk factors at dif-
ferent ecological scales. At the individual level, gender, ethnicity, and work location were
strongly associated with infection risk. At the community level, important predictors of risk
included rural setting, community type, poor access to clean water, close proximity to rivers,
high rainfall in the wettest month, high poverty rate, presence of pigs, and high cattle density.
From a wider perspective, significant geographic variations in risk and the ability to predict
risk based only on environmental and census variables indicate that environmental factors play
a crucial role in driving leptospirosis transmission in Fiji.

The above findings provide an important evidence base to guide the focus of public health
and environmental health interventions at individual, community, and national levels. Health
promotion activities and educational materials should be designed to reach the highest risk
groups including males, farmers, and iTaukei. Public health and environmental health inter-
ventions should target the highest risk communities (villages, rural areas, those in hotspots and
high-risk regions), and include advice on proper management of livestock, avoiding contact
with floodwaters, and minimizing flooding risk (e.g. adequate garbage disposal to reduce the
risk of flooding from blocked streams and drains). At high risk times, e.g. post-flooding, com-
munities should also be reminded about the risk of leptospirosis, the use of protective mea-
sures, and the importance of seeking early medical care if unwell. In smaller communities in
Fiji, where laboratory diagnostic tests are often not available, the predictive risk chart shown in
Fig 8 could assist clinicians with determining the likelihood (pre-test probability) of leptospiro-
sis infection based on a combination of individual-level variables. Broader environmental fac-
tors (both natural and anthropogenic) play a major role in leptospirosis transmission in Fiji,
most of which are beyond the immediate control of individuals or small communities. Effective
environmental health management at the public health and national level will therefore be cru-
cial for the sustainable control of leptospirosis in Fiji and other countries with similar environ-
mental and socio-demographic settings.

Supporting Information
S1 Appendix. Initial 21 pathogenic serovars included in the microscopic agglutination test
(MAT) panels, and the six serovars chosen for the final MAT panel.
(DOCX)
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Author Contributions

Conceived and designed the experiments: CLL CHW SBC MK EJN. Performed the experiments: CLL CHW SBC SJW. Analyzed the data: CLL CHW JHL MCD. Contributed reagents/materials/analysis tools: SBC SJW. Wrote the paper: CLL CHW JHL MCD EJN.

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