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HIV Infection Disrupts the Sympathetic Host–Pathogen Relationship in Human Tuberculosis

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Abstract

The phylogeographic population structure of Mycobacterium tuberculosis suggests local adaptation to sympatric human populations. We hypothesized that HIV infection, which induces immunodeficiency, will alter the sympatric relationship between M. tuberculosis and its human host. To test this hypothesis, we performed a nine-year nation-wide molecular-epidemiological study of HIV-infected and HIV-negative patients with tuberculosis (TB) between 2000 and 2008 in Switzerland. We analyzed 518 TB patients of whom 112 (21.6%) were HIV-infected and 233 (45.0%) were born in Europe. We found that among European-born TB patients, recent transmission was more frequent to be in sympatric compared to allopatric host–pathogen combinations (adjusted odds ratio [OR] 7.5, 95% confidence interval [95% CI] 1.2–infinity, p = 0.03). HIV infection was significantly associated with TB caused by an allopatric (as opposed to sympatric) M. tuberculosis lineage (OR 7.0, 95% CI 2.5–19.1, p < 0.0001). This association remained when adjusting for frequent travelling, contact with foreigners, age, sex, and country of birth (adjusted OR 5.6, 95% CI 1.5–20.8, p = 0.01). Moreover, it became stronger with greater immunosuppression as defined by CD4 T-cell depletion and was not the result of increased social mixing in HIV-infected patients. Our observation was replicated in a second independent panel of 440 M. tuberculosis strains collected during a population-based study in the Canton of Bern between 1991 and 2011. In summary, these findings support a model for TB in which the stable relationship between the human host and its locally adapted M. tuberculosis is disrupted by HIV infection.

Introduction

Host-pathogen co-evolution plays an important role in the biology of infectious diseases [1]. Coevolution between interacting host and pathogen species is difficult to demonstrate formally, but indirect evidence can be obtained by studying geographical patterns, which can indicate local adaptation of particular pathogen variants to geographically matched host variants [2–4]. Local adaptation is often studied using so-called reciprocal transplant experiments, in which the fitness of locally adapted...
Author Summary

Human tuberculosis (TB) caused by Mycobacterium tuberculosis kills 1.5 million people each year. M. tuberculosis has been affecting humans for millennia, suggesting that different strain lineages may be adapted to specific human populations. The combination of a particular strain lineage and its corresponding patient population can be classified as sympatric (e.g. Euro-American lineage in Europeans) or allopatric (e.g. East-Asian lineage in Europeans). We hypothesized that infection with the human immunodeficiency virus (HIV), which impairs the human immune system, will interfere with this host–pathogen relationship. We performed a nation-wide molecular-epidemiological study of HIV–infected and HIV–negative TB patients between 2000 and 2008 in Switzerland. We found that HIV infection was associated with the less adapted allopatric lineages among patients born in Europe, and this was not explained by social or other patient factors such as increased social mixing in HIV–infected individuals. Strikingly, the association between HIV infection and less adapted M. tuberculosis lineages was stronger in patients with more pronounced immunodeficiency. Our observation was replicated in a second independent panel of M. tuberculosis strains collected during a population-based study in the Canton of Bern. In summary, our study provides evidence that the sympatric host–pathogen relationship in TB is disrupted by HIV infection.

(sympatric) pathogen variants is compared to the performance of allopatric pathogen variants [2]. Studies in several invertebrate systems have shown that sympatric pathogens (infection with a phylogeographically concordant strain) tend to outperform allopatric pathogens (infection with a phylogeographically discordant strain) in the corresponding host variants [1, 5–7].

Mycobacterium tuberculosis, the agent causing human tuberculosis (TB) is an obligate human pathogen, which has been affecting humankind for millennia [8–13]. Contrary to previous beliefs linking the origin of TB to animal domestication ~10,000 years ago [14], more recent data suggest that M. tuberculosis evolved as a human pathogen in Africa, and might have co-existed with anatomically modern humans since their origins ~200,000 years ago [8, 10, 12, 13, 15]. Analyses of multiple global strain collections have shown that M. tuberculosis exhibits a phylogeographic population structure consisting of six main phylogenetic lineages associated with different geographic regions and sympatric human populations [9, 11–13, 16–20]. Studies in San Francisco, London, and Montreal have shown that these sympatric host–pathogen associations persist in cosmopolitan settings, even under a presumed degree of host and pathogen intermingling [11, 18, 19]. Moreover, transmission of M. tuberculosis has been shown to occur more frequently in sympatric host–pathogen combinations compared to allopatric host–pathogen combinations [9]. Taken together, these observations are consistent with the notion that the different phylogeographic lineages of M. tuberculosis have adapted to specific sympatric human populations [21].

Based on the assumption that M. tuberculosis has been coevolving with humans, and that M. tuberculosis has locally adapted to sympatric human populations [9], we hypothesized that HIV co-infection will alter this relationship [22]. Specifically, we postulated that because HIV induces immune suppression in humans, and because variation in host immunity will likely play a role in local adaptation, M. tuberculosis strains will cause disease in HIV–infected patients, irrespective of their usual sympatric host–pathogen relationship. To test this hypothesis, we performed a population-based molecular-epidemiological study of HIV–infected and HIV–negative TB patients in Switzerland between 2000 and 2008, a country with a long history of immigration [23].

Results

Patient characteristics and phylogeographic distribution of M. tuberculosis lineages

A total of 518 patients were included in the study, of whom 112 (21.6%) were HIV–infected. Of these 518 patients, 233 (45.0%) were born in Europe (117 in Switzerland), 131 (25.3%) were born in sub-Saharan Africa, 40 (9.3%) in South-East Asia, 36 (7.0%) in the Indian subcontinent, and 24 (4.6%) in Central and South America. Similar to previous studies [9, 10, 19], we found an association between the patient’s place of birth and the particular M. tuberculosis lineages (Figure 1). Lineage 4 (Euro-American lineage) was present in all regions, but particularly common in patients born in Europe and South America. Lineages 5 and 6 (West-African lineages also known as M. africanum [24]) were exclusively found in patients originating from West Africa, whereas Lineage 2 (which includes Beijing) and Lineage 1 were mainly seen in patients originating from the Western Pacific and East Asian regions. Patient characteristics are summarized in Table 1.

Because in European-born patients the host–pathogen combinations defined as sympatric (i.e. Lineage 4/Euro-American lineage in European-born patients) or allopatric (i.e. all other lineages in European-born patients) have been well established [9, 18, 19, 25], we focused on this patient group (n = 233) for the remaining of our analyses.

M. tuberculosis transmission occurs primarily in sympatric host–pathogen combinations

M. tuberculosis transmission was more likely among patients in a sympatric host–pathogen relationship compared to patients in an allopatric host–pathogen relationship (adjusted odds ratio [OR] 7.5, 95% confidence interval [95% CI] 1.2–infinity, p = 0.03, Table 2). Of note, there was no molecular clustering among European-born TB patients infected with an allopatric M. tuberculosis strain. Moreover, we found that only the sympatric Lineage 4 (Euro-American lineage) was detected in European-born clusters as well as in mixed clusters (Table S1), suggesting that sympatric host–pathogen combinations in TB favor transmission.

Impact of HIV infection on the sympatric host–pathogen combination of M. tuberculosis among Europeans

Overall, we found that HIV infection was strongly associated with allopatric M. tuberculosis lineages among European-born TB patients (unadjusted OR 7.0, 95% CI 2.5–19.1, p<0.0001; Table 3). Among the allopatric lineages, we found that Lineages 1, 2 and 3 were more likely to be found in HIV–infected compared to HIV–negative patients when taking the sympatric Lineage 4 (Euro-American lineage) as the reference (Table S2). When investigating the ancestry of the nine HIV–infected patients with an allopatric M. tuberculosis strain, seven patients were confirmed to be of Swiss ancestry over the last three generations, and two patients had Swiss and Italian ancestors in the previous generations (Italian father in the previous generation, or emigrating from Italy in the previous generation).

Several factors could contribute to the association between HIV infection and allopatric lineages. We found that patients with an allopatric M. tuberculosis lineage were younger (median age 41.5 versus 50 years), and had more often a history of frequent
Figure 1. Phylogeography of the six main *Mycobacterium tuberculosis* lineages. A: Phylogenetic tree of the main *M. tuberculosis* lineages described in our study based on the neighbor-joining phylogeny across 23 *M. tuberculosis* complex whole-genome sequences (from Ref. [72]). Numbers on branches refer to the corresponding number of single nucleotide polymorphisms inferred. B: Distribution of the main phylogenetic *M. tuberculosis* lineages among Swiss tuberculosis cases included in the study (n = 518), by geographic origin of the patients. In (A) *Mycobacterium canettii* was used as the outgroup. SNPs, single nucleotide polymorphisms. In (B) the sizes of the pie charts correspond to the number of patients.

**Lineage 1**
(The Philippines and Rim of Indian Ocean)

**Lineage 2**
(East Asia)

**Lineage 3**
(India and East Africa)

**Lineage 4**
(Europe, America and Africa)

**Lineage 5**
(West African 1)

*M. bovis* (animal strain)

**Lineage 6**
(West African 2)

*M. canettii*
(outgroup)
travelling (38.9% versus 4.2%, p < 0.0001). Therefore, we developed a model (Figure 2) to take these and other putative confounding variables into account. These variables included age, sex, country of birth, frequent travelling, contact with the foreign-born population, and non–HIV associated immunosuppression. We considered “TB with an allopatric strain” as the outcome because disease is the only measurable outcome with a sympatric or allopatric M. tuberculosis strain (only diseased individuals can yield a positive mycobacterial culture). Our multivariate analyses revealed that the association between HIV infection and allopatry remained statistically significant after adjustment for all social and patient factors included in our model (OR 5.5, 95% CI 1.5–20.6, p = 0.01, Table 3). Age, sex, being Swiss-born, and non–HIV associated immunosuppression had only a minor effect on the association between HIV infection and TB with an allopatric strain (Table 3). In contrast, a history of repeated travelling to low-income countries had a stronger effect, decreasing the OR to 4.50 (95% CI 1.5–13.6, p = 0.008, Table 3) when adjusting for this variable.

Impact of HIV infection on the sympatric host–pathogen association is a function of the degree of HIV–induced immunosuppression

We also tested if the degree of immunodeficiency as measured by the nadir CD4 T cell count (defined as the lowest CD4 T cell count ever measured in a patient) would have an impact on the association between host population and M. tuberculosis lineage. Among Europeans, the strength of the association between HIV infection and allopatric lineages increased with a decreasing nadir CD4 T cell count in a dose-dependent manner: from an OR of 4.6 (95% CI 0.9–24.7) in patients with a nadir CD4 T cell count of >200 cells/μL to an OR of 12.5 (95% CI 2.6–60.8) in patients with nadir CD4 T cell counts <50 cells/μL (test for trend p < 0.0001; HIV–negative patients as reference). This trend remained statistically significant when adjusting for age, sex, being born in Switzerland, frequent travelling, contact with the foreign-born population, and non–HIV associated immunosuppression (Table 4).

Impact of social mixing on the sympatric host–pathogen association

Increased contact with foreigners originating from high TB burden countries, who have a higher risk of TB [26] and are more likely to have TB caused by an allopatric M. tuberculosis strain, could also lead to an allopatric host–pathogen relationship in European-born patients. However, the association between HIV and allopatry remained statistically significant when adjusting for this variable (Table 3). Furthermore, we examined molecular clusters defined by standard bacterial genotyping [27,28], to test

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>All (n = 233)</th>
<th>Allopatric (n = 18)</th>
<th>Sympatric (n = 215)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, median (IQR), years</td>
<td>49 (36–71)</td>
<td>41.5 (32–45)</td>
<td>50 (37–73)</td>
<td>0.0029</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>126 (54.1)</td>
<td>10 (55.6)</td>
<td>116 (54.0)</td>
<td>0.90</td>
</tr>
<tr>
<td>Origin of birth, n (%)</td>
<td>0.46</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Switzerland</td>
<td>118 (50.6)</td>
<td>11 (61.1)</td>
<td>107 (49.9)</td>
<td></td>
</tr>
<tr>
<td>Europe (without Switzerland)</td>
<td>115 (49.4)</td>
<td>7 (38.9)</td>
<td>108 (50.2)</td>
<td></td>
</tr>
<tr>
<td>Cavitary disease, n (%)</td>
<td>55 (23.6)</td>
<td>4 (22.2)</td>
<td>51 (23.7)</td>
<td>0.99</td>
</tr>
<tr>
<td>Clinical manifestation, n (%)</td>
<td>0.54</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pulmonary</td>
<td>191 (82.0)</td>
<td>16 (88.9)</td>
<td>175 (81.4)</td>
<td></td>
</tr>
<tr>
<td>Extrapulmonary</td>
<td>42 (18.0)</td>
<td>2 (11.1)</td>
<td>40 (18.6)</td>
<td></td>
</tr>
<tr>
<td>Recent TB within families/social surroundings</td>
<td>15 (6.4)</td>
<td>0 (0)</td>
<td>15 (7.0)</td>
<td>0.61</td>
</tr>
<tr>
<td>Frequent travelling</td>
<td>16 (6.9)</td>
<td>7 (38.9)</td>
<td>9 (4.2)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>HIV infection, n (%)</td>
<td>36 (15.5)</td>
<td>9 (50.0)</td>
<td>27 (12.6)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Immunosuppression other than HIV infection, n (%)</td>
<td>24 (10.3)</td>
<td>1 (5.6)</td>
<td>23 (10.7)</td>
<td>0.70</td>
</tr>
<tr>
<td>Most likely source of HIV infection, n (%)</td>
<td>0.94</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heterosexual</td>
<td>15 (41.7)</td>
<td>5 (55.6)</td>
<td>10 (37.0)</td>
<td></td>
</tr>
<tr>
<td>Injecting drug user</td>
<td>9 (25.0)</td>
<td>2 (22.2)</td>
<td>7 (25.9)</td>
<td></td>
</tr>
<tr>
<td>Men having sex with men</td>
<td>7 (19.4)</td>
<td>2 (22.2)</td>
<td>5 (18.5)</td>
<td></td>
</tr>
<tr>
<td>Others/unknown</td>
<td>5 (13.9)</td>
<td>0 (0)</td>
<td>5 (18.5)</td>
<td></td>
</tr>
</tbody>
</table>

1Fisher’s exact test.
2Use of TNF-alpha inhibitors, malignancy, organ transplantation, use of steroids, or methotrexate.
3Among HIV–infected patients (n = 36).
495% CI, 95% confidence interval; IQR, interquartile range.
doi:10.1371/journal.pgen.1003318.t001
the hypothesis that HIV–infected patients were more frequently seen among ethnically mixed clusters where transmission occurred between non-European and European-born cases [29]. We found that the prevalence of HIV infection was similar among TB cases in mixed clusters (5 HIV–infected cases out of 26 cases, 19.2%) and among cases in clusters involving only European-born cases (4 out of 26 cases, 15.4%, see Table S1).

### Sensitivity analyses

When restricting the main analysis (n = 233) to European-born patients without a history of frequent travelling, we found that the association between HIV infection and allopatric TB remained statistically significant (adjusted OR 6.96, 95% CI 1.25–38.88, p = 0.027). Furthermore, we explored associations of socio-demographic and clinical factors with TB with an allopatric strain in a model focusing on HIV–infected European patients only (Figure S1, Table S3): frequent travelling was confirmed to be an important factor, and patients with a low nadir CD4 T cell count tended to be associated with an allopatric TB although the associations did not reach statistical significance (Table S3). Finally, we obtained very similar results for the association between HIV infection and allopatric TB (Table S4), and for the association between the degree of immunodeficiency and allopatric TB (Table S5) when repeating analyses using a Bayesian approach [30], which is more robust when numbers are small.

### Other supporting information

The birth origin of HIV–infected and non-infected patients is shown on a map in Figure S2. The main phylogenetic **M. tuberculosis** lineages stratified by place of birth and HIV status are presented in Table S6.

### Replication in a second panel of **M. tuberculosis** strains

To replicate our main finding, we investigated a second panel of **M. tuberculosis** strains from an ongoing population-based TB study in the Canton of Bern, Switzerland, between 1991 and 2011. Of the 1,642 **M. tuberculosis** isolates analyzed, 1,350 (82.2%) belonged to Lineage 4 (Euro-American lineage), and 292 (17.8%) to non-Euro-American lineages (Lineages 1, 2, 3, 5 or 6). We compared all 40 European-born patients with an allopatric strain (non-Lineage 4) with 400 randomly selected European-born patients with a sympatric strain (Lineage 4). We found that the proportion of HIV infection was 4.5 (95% CI 1.6–11.9) times higher in patients with an allopatric strain compared to patients with a sympatric strain (12.5% versus 2.8%, p = 0.010, Table 5).

### Table 2. Recent transmission of **Mycobacterium tuberculosis** among tuberculosis (TB) cases born in the European region, according to sympatric and allopatric lineages.

<table>
<thead>
<tr>
<th>Lineages</th>
<th>n (%) cases</th>
<th>Association of transmission with lineages</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Clusters</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sympatric</td>
<td>42 (19.5)</td>
<td>173 (80.5)</td>
</tr>
<tr>
<td>Allopatric</td>
<td>0 (0)</td>
<td>18 (100)</td>
</tr>
</tbody>
</table>

Recent transmission was determined by spoligotyping and MIRU-VNTR genotyping which is based on repetitive sequences. Clustered cases were defined as cases belonging to a molecular cluster of TB transmission based on isolates showing an identical genotyping pattern, and unclustered as cases with a unique genotyping pattern. Sympatric was defined as a strain belonging to Lineage 4 (Euro-American lineage), allopatric as a strain belonging to a lineage other than Lineage 4.

1Fisher’s exact test (1-sided). 95% CI, 95% confidence interval; ND, not defined; OR, Odds Ratio. Odds ratios were derived from exact logistic models. Model was adjusted for age group (45 years and younger), being born in Switzerland and recent TB in families or social surroundings. doi:10.1371/journal.pgen.1003318.t002

### Table 3. Unadjusted and adjusted associations between HIV infection and tuberculosis (TB) with an allopatric **Mycobacterium tuberculosis** strain among European patients (n = 233), in the context of other potential factors influencing the risk for an allopatric TB.

<table>
<thead>
<tr>
<th>Variables adjusted for1</th>
<th>OR (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unadjusted</td>
<td>6.96 (2.54–19.08)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Age, sex, Swiss-born</td>
<td>7.54 (2.32–24.55)</td>
<td>0.0010</td>
</tr>
<tr>
<td>Frequent travelling</td>
<td>4.50 (1.49–13.61)</td>
<td>0.0080</td>
</tr>
<tr>
<td>Age, sex, Swiss-Born, frequent travelling, contact with foreign-born population</td>
<td>5.57 (1.49–20.81)</td>
<td>0.011</td>
</tr>
<tr>
<td>Imunosuppression2</td>
<td>7.06 (2.57–19.42)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Age, sex, Swiss-born, frequent travelling, contact with foreign-born population, imunosuppression2</td>
<td>5.51 (1.47–20.61)</td>
<td>0.011</td>
</tr>
</tbody>
</table>

HIV–negative TB patients were used as the reference group.

1See Figure 2 for a graphical overview of associations.

2Imunosuppression other than HIV infection (use of anti-TNF blockers, malignancy, organ transplantation, use of steroids or methotrexate). OR, odds ratio; 95% CI, 95% confidence interval. doi:10.1371/journal.pgen.1003318.t003
Discussion

The phylogeographic distribution of *M. tuberculosis* lineages observed here suggests local adaptation to sympatric human populations. In contrast, we found that allopatric host–pathogen relationships in European-born TB patients were strongly associated with HIV co-infection. The association with HIV infection became stronger in a ‘dose-dependent’ manner in patients with a history of more pronounced immunodeficiency, and was not explained only by frequent travelling to high TB-incidence countries or increased social mixing with the foreign-born population. The association of *M. tuberculosis* lineages with sympatric patient populations reported here is in agreement with previous findings [9,11–13,16–19]. Similarly, our finding that recent TB transmission was more likely to occur in sympatric compared to allopatric host–pathogen combinations supports previous work [9]. Taken together, these data are consistent with local adaptation of *M. tuberculosis* to different human populations, which in turn can be viewed as indirect evidence for coevolution between *M. tuberculosis* and its human host [1–4,9–13].

**Table 4.** Association between the degree of immunodeficiency and tuberculosis with an allopatric *Mycobacterium tuberculosis* strain among European patients (n = 233).

<table>
<thead>
<tr>
<th>Degree of immunodeficiency</th>
<th>n</th>
<th>Unadjusted</th>
<th></th>
<th>Adjusted</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>OR</td>
<td>95% OR</td>
<td><em>P</em> value</td>
<td>OR</td>
</tr>
<tr>
<td>Nadir CD4 T cell count (CD4 cells/μl)</td>
<td></td>
<td>&lt;0.0001</td>
<td></td>
<td></td>
<td>0.0050</td>
</tr>
<tr>
<td>HIV–negative</td>
<td>197</td>
<td>1.0</td>
<td>(ref)</td>
<td>(ref)</td>
<td>1.0</td>
</tr>
<tr>
<td>≥200</td>
<td>11</td>
<td>4.64</td>
<td>(0.87–24.70)</td>
<td>2.56</td>
<td>(0.37–17.65)</td>
</tr>
<tr>
<td>50–199</td>
<td>17</td>
<td>6.43</td>
<td>(1.74–23.70)</td>
<td>6.96</td>
<td>(1.18–41.11)</td>
</tr>
<tr>
<td>&lt;50</td>
<td>8</td>
<td>12.53</td>
<td>(2.58–60.84)</td>
<td>13.0</td>
<td>(1.54–109.75)</td>
</tr>
</tbody>
</table>

Model was adjusted for age, sex, Swiss-born, frequent travelling, contact with foreign-born population, and immunosuppression other than HIV infection (see Figure 2 for a graphical overview).

*P* values of linear tests for trend are shown.
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We found that TB allopatric host-pathogen combinations were strongly associated with HIV infection in a nation-wide study and a second panel of strains from one Canton of Switzerland. This supports the notion that *M. tuberculosis* lineages have evolved subtle differences in their interaction with different human immune systems. However, in the presence of HIV-induced immunodeficiency, any *M. tuberculosis* lineage seems to cause disease in a given human host. *M. tuberculosis* is an obligate human pathogen which lives in constant interaction with the host immune system [31]. Human populations, however, are known to differ genetically and immunologically [15]. The clinical disease reflects host-dependent immune-pathological processes [31]. In other words, while initially triggered by the pathogen, it is the host immune response which is ultimately responsible for the chronic inflammation and associated tissue destructions. These processes contribute to the successful transmission of *M. tuberculosis* [22,32]. On the other hand, only 5–10% of the 2 billion individuals estimated to be latently infected with *M. tuberculosis* globally will develop active TB during their lifetime [33–35]. Hence most of the time, humans are able to control the infection. In our study, we chose culture-confirmed TB lifetime [33–35]. Hence most of the time, humans are able to control the infection. In our study, we chose culture-confirmed TB.

### Table 5. HIV status in tuberculosis (TB) patients with an allopatric compared to patients with a sympatric *Mycobacterium tuberculosis* strain among European patients in a second panel.

<table>
<thead>
<tr>
<th>HIV status</th>
<th>TB cases, n (%)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All (n = 440)</td>
<td>Allopatric (n = 40)</td>
</tr>
<tr>
<td>HIV–infected</td>
<td>16 (3.6)</td>
<td>5 (12.5)</td>
</tr>
<tr>
<td>HIV–negative</td>
<td>424 (96.4)</td>
<td>35 (87.5)</td>
</tr>
</tbody>
</table>

Patient isolates were obtained from a population-based TB study (n = 1,642) in the Canton of Bern, Switzerland, diagnosed between 1991 and 2011. Sympatric was defined as a strain belonging to Lineage 4 (Euro-American lineage), allopatric as a strain belonging to a lineage other than Lineage 4. Fisher’s exact test. 95% CI, 95% confidence interval.

doi:10.1371/journal.pgen.1003318.t005

In conclusion, our data suggest that the phylogeographical host-pathogen relationship in human TB is particularly frequent in native populations of this region [53]. Similarly, a study of the bacterial genome evolution of an asymptomatic *Escherichia coli* bacteriuria strain showed adaptation at the genomic level in distinct human hosts [54]. No similar experimental work has yet been carried out in TB. However, several studies have reported associations between human genetic polymorphisms and particular *M. tuberculosis* lineages [55–59], indicating possible interaction between human and *M. tuberculosis* variation. Whether such variation in pathogen and host genetics can be attributed to co-evolution will be difficult to demonstrate conclusively, but the data presented here support this possibility.

The strength of our study was that we used a nation-wide sample to specifically look at the impact of HIV infection on the host-pathogen relationship in human TB. Yet, our study is limited by the relatively small sample size, and the difficulty to quantify the complex social context through which the host-pathogen relationship is influenced in human TB. In addition, we looked at European-born patients only, because sympatric and allopatric host-pathogen combinations are more easily defined for this patient population [9,18,19,25]. Additional studies in large cosmopolitan cities of Asia and Africa would be required to test whether the association between allopatric TB and HIV holds true in these settings. Ultimately, detailed experimental work is needed to establish the biological basis of the host-pathogen association in human TB.

In conclusion, our data suggest that the phylogeographical host-pathogen relationship in TB influences transmission patterns. Among the studied European-born TB patients, we showed that HIV infection disrupts the sympatric host-pathogen relationship in human TB, and that this effect increased as a function of immunodeficiency. Various interactions between HIV and *M. tuberculosis* at the cellular level make an association biologically plausible [42,43]. Further studies are needed to investigate the impact of HIV on the genetic population structure of *M. tuberculosis* with its consequences for transmission and clinical manifestations in high TB burden countries [36]. This will lead to a better understanding of biological factors that shape the current HIV/TB syndemic [60].

## Methods

### Study setting

The Swiss Molecular Epidemiology of Tuberculosis (SMET) study is a collaborative project between the Swiss HIV Cohort...
Study (SHCS), the National Center for Mycobacteria, diagnostic microbiology laboratories, departments of respiratory medicine and public health, and the Federal Office of Public Health (FOPH) [29,61,62]. The overarching aims were to examine the genetic population structure of M. tuberculosis and the associations between strain variation, patient origin, and clinical characteristics in HIV–infected and HIV–negative TB patients in Switzerland. Further information on the SMET project is available at www.tb-network.ch. All participating sites are listed in the Acknowledgements.

The SHCS is a prospective observational study of HIV–infected individuals followed up in HIV outpatient clinics in Switzerland [63]. All HIV–infected patients diagnosed with TB between 2000 and 2008 whose M. tuberculosis complex (MTBC) isolate was available were included in the SMET study [29]. Furthermore, we randomly selected 288 from the 4,221 culture-confirmed TB cases reported to the National TB Surveillance Registry during the same period (approximately three cases for one HIV–infected TB case within the SHCS). Finally, all reported drug-resistant TB cases were included. Two M. bovis isolates were excluded from this study as they are animal-adapted species within the MTBC and therefore represent a different host–pathogen relationship.

Clinical data collection and definitions

We obtained clinical data by standardized questionnaires sent to the treating physicians and extracted relevant data from the SHCS database. We collected socio-demographic data (age, sex, origin of birth, citizenship, legal status, immunosuppressive therapy, risk factors for TB such as recent TB within family or immediate social surroundings in the last two years), laboratory parameters (CD4 cell count and plasma HIV RNA in HIV–infected cases) and clinical information (site of disease, radiography findings). Chest radiography parameters were consolidation, cavitations, enlarged intrathoracic lymph nodes and pleural thickening. Any drug resistance was defined as any resistance to isoniazid, rifampicin or ethambutol as reported to the FOPH. Most TB cases in Switzerland are treated under the guidance of experienced treating physicians and, in turn, have a direct effect on the outcome variable. We included age and sex in our model as risk factors for infection and disease [37]. We also considered contact with the foreign-born population who have a higher risk for TB compared to the native Swiss population [26] and who have a higher risk of exposure to “foreign” M. tuberculosis strains. Finally, we included frequent traveling to countries with a high TB burden, which increases exposure risk and thus potentially infection risk with “foreign” M. tuberculosis strains (Figure 2).

Statistical analyses

We used χ² tests or Fisher’s exact tests to assess differences between groups in binary variables, and the Wilcoxon rank sum test for continuous variables (Table 1, Table 2). Univariate and multivariate exact logistic regression models were fitted to estimate the association between transmission as defined by molecular clustering and patients with sympatric M. tuberculosis lineages (patients with allopatic lineages were used as the reference, Table 2). Results were presented as ORs unadjusted and adjusted for age group, being born in Switzerland and recent TB in families or social surroundings. To assess the association of HIV infection with allopatic TB, we fitted univariate and multivariate logistic models (Table 3), and presented ORs unadjusted and adjusted for age, sex, Swiss-born, frequent travelling, contact with foreign born populations, and/or immunosuppression. We used univariate and multivariate logistic models to estimate the association between the degree of immunodeficiency and allopatic TB (Table 4), and presented ORs unadjusted and adjusted for age, sex, Swiss-born, frequent travelling, contact with foreign-born populations, and immunosuppression other than HIV infection. Finally, we determined statistical significance of HIV prevalence in patients with allopatic M. tuberculosis lineages compared to patients with sympatric lineages using Fisher’s exact tests (Table 5). All analyses were performed in Stata version 11.1 (Stata Corporation, College Station, TX, USA).

Molecular analyses

Mycobacterial isolates were cultured and DNA extracted according to standard laboratory procedures. We used spacer oligonucleotide typing (spoligotyping) and 24-loci mycobacterial interspersed repetitive units (MIRU-VNTR) which are based on repetitive DNA sequences as genotyping tools with high discriminatory power to identify recent TB transmission [29,64–66]. Data were analyzed with the MIRU-VNTRplus online tool (http://www.miru-vntrplus.org). Molecular clusters were defined as a group of completely identical isolates in the spoligotyping and MIRU-VNTR pattern indicating a chain of TB transmission. In addition, we used single nucleotide polymorphisms (SNPs) as stable genetic markers to define the main phylogenetic M. tuberculosis lineages [67]. Lineages were determined by SNPs using multiplex real-time PCR with fluorescence-labeled probes (Taqman, Applied Biosystems, USA) adopted from previous studies [9,12,67,68]. The SNP used to define Lineage 4 was originally described by Sreevatsan et al. [69] and shown to be specific to this lineage [9].

Graphical models

Graphical models were built using the principles of directed acyclic graphs [70]. Our model considered infection and disease as a combined outcome (“TB with an allopatic strain”). Our hypothesis that HIV infection causes TB with an allopatic strain is shown as a potentially causal direct effect, and risk factors potentially influencing this effect are shown in the hypothetical direction. Mediators represent variables that are caused by the independent variable and, in turn, have a direct effect on the outcome variable. We included age and sex in our model as risk factors for infection and disease [37]. We also considered contact with the foreign-born population who have a higher risk for TB compared to the native Swiss population [26] and who have a higher risk of exposure to “foreign” M. tuberculosis strains. Finally, we included frequent traveling to countries with a high TB burden, which increases exposure risk and thus potentially infection risk with “foreign” M. tuberculosis strains (Figure 2).
distributions. Bayesian statistics are less sensitive to errors when calculating estimators and CIs in small datasets.

Second panel of \textit{M. tuberculosis} strains

We obtained 1,642 \textit{M. tuberculosis} isolates from all TB cases (n = 1,940, 84.6\%) notified in the Canton of Bern, Switzerland, between 1991 and 2011. For all patient isolates, we determined the main phylogenetic \textit{M. tuberculosis} lineages. Of these, we included all patient isolates belonging to a non-Euro-American lineage (Lineage 1, 2, 3, 5 or 6) from European-born TB patients (40 of a total of 292 isolates belonging to lineages other than Lineage 4). Furthermore, we randomly selected control strains belonging to a total of 292 isolates belonging to Lineage 4. European ancestry was confirmed in HIV–infected patients with an allopatic \textit{M. tuberculosis} strain. Finally, we determined the HIV status in these patients using the same procedures as in the main sample.

Ethics approval

The study was approved by the ethics committee of the Canton of Bern, Switzerland. Written informed consent was obtained from all patients enrolled in the SHCS. For patients outside the SHCS, written informed consent was obtained by the treating physicians. In some cases informed consent could not be obtained from the patient because he or she could not be located or was known to have died. For these cases we obtained permission from the Federal expert commission on confidentiality in medical research to use the data provided by the treating physician.

Supporting Information

Figure S1 Graphical model showing direct potential effects on tuberculosis (TB) with an allopatic \textit{Mycobacterium tuberculosis} strain among HIV–infected patients.

Figure S2 Distribution of tuberculosis (TB) cases included in the study, by origin of birth and HIV status.

Table S1 Distribution of the main phylogenetic \textit{Mycobacterium tuberculosis} lineages according to molecular clusters involving either non-European-born tuberculosis (TB) cases only, European-born cases, or mixed clusters involving both non-European-born and European-born TB cases.

Table S2 Crude and adjusted analysis comparing HIV–infected and HIV–negative tuberculosis (TB) patients born in Europe (n = 233) across the four most frequent \textit{Mycobacterium tuberculosis} lineages.

Table S3 Associations of socio-demographic and clinical factors with tuberculosis (TB) with an allopatic \textit{Mycobacterium tuberculosis} strain among HIV–infected European patients (n = 36).

Table S4 Associations between HIV infection and tuberculosis (TB) with an allopatic \textit{Mycobacterium tuberculosis} strain among European patients (n = 233) in the context of other potential factors influencing this association, using Bayesian statistics and presented as unadjusted or adjusted odds ratios.

Table S5 Association between the degree of immunodeficiency and tuberculosis (TB) with an allopatic \textit{Mycobacterium tuberculosis} among European patients (n = 233) using Bayesian statistics.

Table S6 Comparing the main phylogenetic \textit{Mycobacterium tuberculosis} lineages, by HIV status and birth region.

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\textbf{Swiss HIV Cohort Study}


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References


