Importance of Cough and *M. tuberculosis* Strain Type as Risks for Increased Transmission within Households

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**Abstract**

**Rationale:** The degree to which tuberculosis (TB) is transmitted between persons is variable. Identifying the factors that contribute to transmission could provide new opportunities for TB control. Transmission is influenced by host, bacterial and environmental factors. However, distinguishing their individual effects is problematic because measures of disease severity are tightly correlated, and assessing the virulence of *Mycobacterium tuberculosis* isolates is complicated by epidemiological and clinical confounders.

**Objectives:** To overcome these problems, we investigated factors potentially associated with TB transmission within households.

**Methods:** We evaluated patients with smear-positive (≥2+), pulmonary TB and classified *M. tuberculosis* into single nucleotide polymorphism genetic cluster groups (SCG). We recorded index case, household contact, and environmental characteristics and tested contacts with tuberculin skin test (TST) and interferon-gamma release assay. Households were classified as high (≥70% of contacts with TST ≥10 mm) and low (≤40%) transmission. We used logistic regression to determine independent predictors.

**Result:** From March 2008 to June 2012, we screened 293 TB patients to enroll 124 index cases and their 731 contacts. There were 23 low and 73 high transmission households. Index case factors associated with high transmission were severity of cough as measured by a visual analog cough scale (VACS) and the Leicester Cough Questionnaire (LCQ), and cavitation on chest radiograph. SCG 3b strains tended to be more prevalent in low (27.3%) than in high (12.5%) transmission households (p = 0.11). In adjusted models, only VACS (p = 0.001) remained significant. SCG was associated with bilateral disease on chest radiograph (p = 0.002) and marginally associated with LCQ sores (p = 0.058), with group 3b patients having weaker cough.

**Conclusions:** We found differential transmission among otherwise clinically similar patients with advanced TB disease. We propose that distinct strains may cause differing patterns of cough strength and cavitation in the host leading to diverging infectiousness. Larger studies are needed to verify this hypothesis.

**Introduction**

Tuberculosis (TB) remains one of the most important threats to global health. Although TB is the prototypic disease with airborne transmission [1], there is both experimental [2,3] and epidemiologic [4,5,6,7,8,9] evidence of marked variability in transmission of *M. tuberculosis* (MTB) from patients with pulmonary TB. The reasons underlying this variability remain poorly understood in part because of the inherent complexity of studying a biological phenomenon resulting from a complex web of interactions between two human hosts—the source case and the exposed contact—and the infecting pathogen within an array of environments. Yet, these differences have important implications for vaccine development and global TB control.
Most studies of transmission and disease in household contacts have focused on host and environmental factors [7,10]. These studies fail to explain differences in epidemiologic behavior of clinical isolates of MTB reported in outbreaks. In fact, there is a growing body of experimental evidence suggesting that MTB strains have variable degrees of virulence [11,12,13,14,15,16,17,18,19]. Although a great deal is known about the genome and proteome of MTB, the genetic basis for most of the cited differences in strain behavior is unknown in most cases [20]. Despite this growing body of literature, there have been no systematic studies designed to investigate the role of strain difference as a determinant of the critical epidemiologic and clinical parameters such as transmissibility, resistance to infection and progression to disease. Importantly, most of the cited studies investigating the virulence potential of MTB isolates have used banked clinical isolates with limited knowledge about the epidemiologic and clinical conditions present at the time the isolates were collected [11,12,13,14,15,16,17,18,19,20].

We conducted a household contact study in Vitória, Brazil to test the hypothesis that genetic strain variation in MTB was a determinant of the extent of transmission of infection in household contacts of smear-positive cases of pulmonary TB. We sought to determine whether some clinical strains of MTB were more transmissible than others, while controlling for other host and environmental factors involved in transmission.

**Methods**

**Ethics Statement**

The study was approved by the Institutional Review Boards of Boston University Medical Campus and New Jersey Medical School – Rutgers University (formerly University of Medicine and Dentistry of New Jersey), the Comitê de Ética em Pesquisa do Centro de Ciências da Saúde - Universidade Federal do Espírito Santo and the Comissão Nacional de Ética em Pesquisa (CONEP). We obtained written informed consent and assent in Portuguese in accordance with age-specific ethical guidelines of participating institutions.

**Study population**

This study was conducted at the The Núcleo de Doenças Infecciosas (NDI) located in Vitória, the capital city of the State of Espírito Santo, Brazil. The NDI has organized a network of five laboratories in the metropolitan region of Vitória that serve 16 TB clinics distributed in the municipalities of Vitória, Cariacica, Serra and Vila Velha. With approximately 1,400 cases/year, the annual TB incidence in Espírito Santo is 38/100,000 inhabitants. The prevalence of HIV infection in Espírito Santo is <1% in the general population, and 7% in TB cases [21].

All consecutive pulmonary TB patients identified through the NDI clinic network were eligible to participate in this study, provided they fulfilled the following inclusion criteria: 1) age ≥18 years with cough ≥3 weeks; 2) new TB episode with ≥1 sputum specimen with acid-fast bacilli (AFB) ≥2+ with subsequent MTB growth in culture; and 3) living with ≥3 household contacts. We excluded index cases who were HIV-infected (or refused HIV testing), had a history of TB treatment, or who were too ill to consent, unable to understand, or to comply with the study protocol. To minimize differences in exposure time between study households, index TB cases were screened and enrolled within the first 2 weeks after they first presented to the municipal TB clinic. A household contact was defined as an individual of any age fulfilling at least one of the following criteria of close contact with the index TB cases for ≥3 months before enrollment: 1) sleeping under the same roof ≥5 days/week; 2) sharing meals ≥5 days/week; 3) watching TV nights or weekends, and; 4) other significant contact (85% of these visited the household >18 days/month).

**Measurements**

**TB cases.** We collected demographic and clinical information about TB patients and recorded their cough severity using two measurements: i) a self-reported visual analog cough scale (VACS) [22], and ii) the Leicester Cough Questionnaire (LCQ) [23,24]. The LCQ is a 19-item, patient-derived questionnaire that measures the physical, psychological and social effects of chronic cough in adults; the final score (range 3–21) is inversely related to cough severity (e.g. higher scores indicate weaker cough). Because of the inherent complexity of the LCQ, study staff administered the LCQ only to TB patients deemed reliable historians (over 95%). We obtained up to three sputa specimens for AFB smear microscopy (auramine O fluorescent stain) [25] and culture (Ogawa-Kudoh method). MTB isolates underwent single nucleotide polymorphism (SNP) analysis as described previously [26] and were categorized into one of nine SNP genetic cluster groups and subgroups (SCG), as described previously [27]. The radiological extent of lung disease was graded on a four-category ordinal scale (normal, minimal, moderate and far-advanced) by an experienced physician [28]. All patients were offered standard TB treatment according to Brazilian guidelines [29,30]. Study staff visited the participants’ dwellings to verify the identity of each contact, measure individual contact time, and to perform an environmental evaluation (crowding and ventilation).

**Household contacts.** We followed recommendations of the Brazilian TB Control Program for household contact investigations [29,30]. We recorded demographic and clinical characteristics of contacts, and evaluated them for MTB infection with both TST and IGRA (Quantiferon Gold-In-Tube, Qiagen). To ensure accurate TST readings, only staff trained by the National TB Program were responsible for TST application, and we completed inter- and intra-reader evaluations (kappa >90%) prior to opening study enrollment. Briefly, we placed two units of R23 (SSI, Denmark) provided by the Brazilian Ministry of Health on the forearm of contacts using the Mantoux method; the diameter of induration was measured in millimeters between 72 and 96 hours. Household contacts with a TST<10 mm at baseline were re-tested after 8–12 weeks to identify TST conversion, using two different criteria for conversion: **Criterion 1 (Brazilian Guidelines):** 1st TST<10 mm, 2nd TST≥10 mm, and difference between 1st and 2nd TST≥10 mm. **Criterion 2:** 1st TST<5 mm, 2nd TST≥10 mm, and difference between 1st and 2nd TST≥6 mm [31]. At 8–12 weeks, blood for IGRA testing was obtained before TST placement to reduce TST-induced IGRA boosting [32]. Following Brazilian recommendations [29], we referred contacts with TST≥10 mm and secondary TB suspects for evaluation and treatment by the corresponding Municipal TB Clinic.

**Household transmission categories**

We categorized household transmission categories according to the percentage of contacts in each household that had a TST≥10 mm by 8–12 weeks: low (LT) transmission (≤40%) or high (HT) transmission (≥70%). This definition includes contacts with TST≥10 mm at either baseline or week 8–12, providing a more complete measure of the total infection burden for each household. Originally, 25% and 75% cutoffs for LT and HT were planned but relaxed to 40% and 70%, respectively, based solely on increasing the numbers in the two categories; no other factors were examined when the cutoffs were revised. A group classified as intermediate transmission (IT) households (41–69%) contacts with
purposes of the determination of the transmission category. Reverse; these two contacts were considered TST-positive for determined to have been infected by the index case and not the with co-prevalent TB disease and based on symptom history were determined the transmission category; their inclusion or exclusion had little effect on the category (data not shown). Two contacts were diagnosed on the category (data not shown). Two contacts were diagnosed as control households.

**Statistical methods**

Because this study used a household contact design, characteristics are at both a household level (including data about the index case) and at an individual contact level, nested within households. Households were classified into three phenotypes, i.e., low, intermediate, and high transmission. Comparisons were made between LT and HT households, and also across the three phenotypes. Unadjusted associations of household level data to transmission phenotype assumed independence across households. Logistic regression was used to obtain odds ratios of household level characteristics, but in the case of sparse data, exact logistic regression and corresponding odds ratios and tests were substituted [33]. Characteristics from individual contacts within a household were potentially correlated and, therefore, were evaluated using a generalized estimating equation (GEE) approach with an independent working correlation structure. We assumed a binomial distribution with a logit link when comparing LT and HT households and a multinomial distribution with a cumulative logit link when comparing LT, IT, and HT households. Multiple regression modeling focused on comparing LT vs. HT households and used the GEE approach to account for within household correlation. An adhoc approach was used to determine characteristics for the final multiple regression model. An automatic stepwise selection procedure was used to select household level characteristics, and then individual contact level covariates were candidate for addition to the model in a manual stepwise fashion. SCG and contact age were then added to the model. Interactions of SCG with measures of disease severity used simplified models because of estimation problems. Covariates of questionable reliability were excluded from consideration. Because of their strong negative correlation (Pearson correlation \(-0.72\)) VACS and LCQ were not candidates for inclusion in a single multivariable model. Direct comparison of VACS and LCQ showed that VACS was a superior predictor of transmission category based on both Akaike Information criterion (AIC) and likelihood score statistic. Moreover, LCQ is time consuming and difficult to administer and score. Therefore, multivariable analyses only considered inclusion of VACS. All analyses were conducted in SAS 9.2 and testing is two-sided.

**Results**

Between February 2008 and June 2012, we identified 1,071 potentially eligible TB patients, 743 were determined ineligible and not referred to the study, 35 refused study participation, and 743 were determined ineligible (71.8%), and 4 or 6a (90.0%) but these differences were not statistically significant (p = 0.060). Similarly, cavitations were present in SCG 3b (65.0%), 5 (72.7%) (p = 0.002), and less likely to have infiltrates in other SCG (72.7%) (p = 0.002), and less likely to have bilateral disease (15.0%), vs. SCG 5 (51.2%) and 3b (90.0%) but these differences were not statistically significant (p = 0.039).

**Index TB patients and dwellings**

The median age of TB cases was 36 years (range 25, 46), 64% were male, and most had advanced TB disease as measured by duration of cough (median 12 weeks), sputum AFB smear (82% AFB 3+), or extent of lung disease on chest radiograph (Table 1). The severity of cough (as measured by VACS and LCQ) and the presence of cavitation on chest radiograph in index TB cases increased as the proportion of household members with MTB infection increased. There were no differences in socio-economic variables (marital status, highest education level, occupation and average salary) between household transmission categories, except that families living in HT households were more likely to rent (P = 0.012). When compared to LT households, contacts from IT and HT families were more likely to self-report close contact with another person with TB other than the index case (P = 0.02).

**Genotypic strain analysis**

The two most prevalent strains among TB patients were group 5 (73.9%) and group 3b (16.8%) and their distribution across the three household transmission categories differed qualitatively, although not statistically significantly (omnibus P = 0.38) (Table 1). Group 3b strains were more prevalent in LT (27.3%) and IT (20.0%) households compared to HT (12.5%) households; conversely, group 5 strains were most prevalent in HT (76.4%) households compared to LT (68.2%) and HT (72.0%) households. The median [interquartile range (IQR)] LCQ score was 12 (7, 21) for patients infected with SCG 3b strains, 11 (6, 21) for SCG 5, and 10 (5, 16) for other SCG groups combined (p = 0.056), trending toward less coughing in patients infected with a SCG 3b isolate. Similarly, the median [IQR] VACS was 6 (0, 10) for SCG 3b TB patients (N = 19), 7 (0, 10) for SCG 5 (N = 83), and 9 (3, 10) for all other groups (N = 11) combined (p = 0.11), indicating patients infected with a SCG 3b strain had a trend toward a weaker cough (Figure 2). Patients with SCG 3b isolates were less likely to have bilateral disease (15.0%), vs. SCG 5 (31.2%) and other SCG (72.7%) (p = 0.002), and less likely to have infiltrates in the upper lung: SCG 3b (80.0%), 5 (95.3%), and 4 or 6a (100.0%), with the differences being marginally significant (p = 0.060). Similarly, cavitations were present in SCG 3b (65.0%), 5 (71.8%), and 4 or 6a (90.0%) but these differences were not statistically significant (p = 0.39).

**Household contacts**

This analysis includes a total of 731 household contacts that fulfilled the study definition of close and prolonged contact time with the index case (categorized by the first definition met): 63% slept under the same roof \( \pm 5 \) days/week, 12% watched TV on nights/weekends, 11% shared meals \( \pm 5 \) days/week and 14% had other significant contact (85% visited the household \( \geq 18 \) days/month). Most (90%) households were evaluated within \( \pm 14 \) days of the index TB cases starting anti-tuberculous treatment. Household contacts were young (median age 20 years), but covering a broad age range (3 months to 87 years); 56.3% were female; 77.0% had a BCG vaccination scar; 99% had no prior diagnosis of MTB and 98% reported no HIV-infection (Table 1). Enrolled contacts spent considerable time in close contact with their family member with TB. Contacts were comparable across transmission categories in terms of socio-economic condition (marital status, education level, occupation and salary), family composition (i.e. no new contacts left the household or died in 3
months prior to study inception), alcohol and tobacco consumption, and history of co-morbidities.

At baseline, TST was positive in 57.3% of contacts and the distribution varied according to transmission category (by design) and contact age (Table 1). Whereas the percentage of contacts with TST ≥ 10 mm living in LT and IT households increased with age, it was similar across age strata in HT households. Overall, the frequency of IGRA positivity at 8–12 weeks was approximately 15% lower than TST but showed a similar age-dependent increase in prevalence of infection. The median TST diameter in contacts with TST ≥ 10 mm (1st or 2nd TST) increased in step with household transmission categories (P < 0.001).

Household transmission categories

Transmission categories of households changed considerably between baseline and week 8–12 as a result of TST conversion (Tables 2 and 3). At baseline, 41, 30 and 53 of the sampled households were LT, IT and HT, respectively; by 8–12 weeks, 43% of LT and IT household progressed and only 23 remained as LT households (Table 2). The use of two different criteria to define TST conversion identified the same number of contacts with newly acquired infection, however the same contacts were not identified by both criteria (Table 3); of the 56 contacts that underwent TST conversion by either criterion, 44 (79%) fulfilled both TST conversion criteria. Most TST conversions occurred in households initially classified as low or intermediate households, as
| Characteristic | Final Household transmission category | P value (Multinomial) | P value (LT vs. HT) |
|---------------|---------------------------------------|-----------------------|---------------------|
| **Index Cases** | | | |
| **Age (years)** | Total = 124 | Low (LT) = 23 | Intermediate = 28 | High (HT) = 73 | 0.34K | 0.47W |
| **Sex** | | | | | 0.77C | 0.58C |
| **Cough visual analog scale** | | | | | 0.003K | 0.002W |
| **Leicester cough questionnaire** | | | | | 0.007K | 0.007W |
| **Weeks sick prior to enrollment** | | | | | 0.26K | 0.57W |
| **Housing** | | | | | 0.017T | 0.11E |
| **Sputum AFB smear (highest grade)** | | | | | 0.49C | 0.30C |
| **Extent of lung disease on chest radiograph** | | | | | 0.20J | 0.24T |
| **Cavitations** | | | | | 0.012T | 0.023T |
| **SNP cluster group** | | | | | 0.38T | 0.46E |
| **Number of contacts enrolled in study** | | | | | 0.10K | 0.98W |
| **Household Contacts** | | | | | 0.85S | 0.40S |
| **Age (years)** | Total = 731 | Low (LT) = 132 | Intermediate = 183 | High (HT) = 416 | 0.65S | 0.80S |
| **Sex** | | | | | 0.85S | 0.40S |
| **BCG scar** | | | | | 0.72S | 0.86S |
| **Close contact with person with TB other than index case** | | | | | 0.020S | <0.001S |
| **TST and IGRA status, n/N (%)** | | | | | | |
| **1st TST 10 mm** | 407/710 (57.3) | 17/132 (12.9) | 78/180 (43.3) | 312/398 (78.4) | | |
| **1st or 2nd TST 10 mm** | 488/691 (71.7) | 27/127 (21.3) | 103/174 (59.2) | 358/380 (94.2) | | |
| **≤5 years** | 48/85 (56.5) | 0/20 (0.0) | 9/23 (39.1) | 39/42 (92.9) | | |
Factors associated with high MTB transmission

We analyzed index TB patient, household contact and environmental factors associated with HT households (Table 4). In unadjusted analyses, factors associated with HT were the presence of cavitation in chest radiograph (Odds Ratio (OR) 3.13, 95% confidence interval (CI): 1.13, 8.69; P = 0.028) and stronger cough as measured by VACS (OR 1.36 per unit increase, 95% CI: 1.13, 1.67; P = 0.001) or LCQ (OR 1.24 per unit decrease, 95% CI: 1.07, 1.45; P = 0.005) in index TB cases. A group 5 SCG MTB isolate had more than twice the estimated odds of being associated with HT compared to group 3b, although it was not statistically significant (OR 2.44, 95% CI: 0.72, 7.93; P = 0.14). In a logistic regression model adjusted for other covariates with a GEE approach for estimation, the only variable that remained independently associated with HT was severity of cough in the index TB case (VACS adjusted OR per unit increase 1.53, 95% CI: 1.07, 2.19; P = 0.019). The presence of cavitation on chest radiograph trended toward significance (P = 0.078), age of the contact (P = 0.17) and SCG (p = 0.40) were not significantly associated with a HT household. We found no evidence of interaction between SCG and either cavitation (p = 0.50) or VACS (p = 0.49) but were underpowered for such comparisons and in fact had estimation problems.

Discussion

In this household contact study of smear-positive, culture confirmed pulmonary TB patients in Brazil, we systematically identified households with low and high transmission of MTB during a four-year period. Our study design and use of both TST and IGRA to measure prevalent and incident infection in exposed contacts showed heterogeneity of MTB transmission within households. We also found that cough severity in the index case is significantly associated with high transmission and that the presence of cavitation on chest radiograph is also associated, albeit marginally, with increased transmission. Our findings indicate that the infecting strain may result in distinct patterns of pathophysiology of disease in index cases leading to differences in their infectiousness.

The importance of cough frequency as a surrogate for TB infectiousness is well recognized [34,35,36,37]. Overnight cough frequency has been associated with the proportion of contacts with a positive TST [35]. However, the objective evaluation of cough, an inherently subjective symptom, is challenging and we are unaware of previously published data relating cough strength or other measures of cough physiology with MTB transmission. Also, hosts may be more infectious during a specific time of day so there is a need for an objective and quantitative measure of infectiousness [35,38]. Similarly, a hallmark of TB disease is the formation of lung cavitation, the result of liquefaction of tissue after the initial granulomatous response [39]. With progression of disease, cavitation becomes more common, occurring in 45–91% of cases [40]; when present, cavities are associated with a higher bacillary load and increased transmission [35,39,41] when compared to patients with nodular lesions in the lung parenchyma [42,43,44]. However, measures of pulmonary TB disease severity such as cough strength, cavitation, and bacterial load in sputum are tightly correlated and distinguishing their individual effects on MTB transmission can be challenging. We addressed this in our study design by using two separate instruments to measure cough (VACS and LCQ), recording the presence of cavitations on chest radiograph and restricting enrollment to smear positive (≥2+), culture-confirmed TB patients in an effort to maximize and homogenize already proven exposure across households. Whereas the VACS and LCQ have been validated in other conditions with pulmonary TB [45,46], the VACS and LCQ have been validated in other conditions with pulmonary TB [45,46].
based on phenotypic properties in varying conditions such as increased production of pro-inflammatory cytokines in the source or contact cases, growth characteristics in liquid cultures, macrophages, and THP1 cells, and growth kinetics in animal experiments [16,17,18]. Although some of the in-vitro data show conflicting results [45,46], current thinking is that virulent strains subvert the immune system by causing a reduced inflammatory response, which in turn results in impaired bacterial control by the host, more rapidly progressive disease, and enhanced transmission. Strain virulence also can be inferred from the capacity of MTB clones to spread in human populations, as measured by the identification of strain clusters using genotyping techniques [11,15,19,47]. However, it remains unclear whether there is a microbial basis to explain why some clinical isolates cause widespread disease and other closely related strains remain limited in spread [46,48,49]. Our study suggests that SCG 3b strains are associated with diminished transmission but that this may be due to weaker cough and decreased cavity formation, opening the possibility of strain-dependent variable lung parenchymal pathophysiology and/or inflammation in the index TB patient. In fact, the hallmark of the local immune response in pulmonary TB is an unregulated activation of cells and expression of pro-inflammatory and anti-inflammatory cytokines and chemokines [50]. SCG 3b roughly corresponds to the H spoligotype; SCG5, the other SCG which was commonly observed in our index cases, roughly corresponds to the LAM spoligotype. Both LAM and H spoligotypes are members of the Euro-American LSP/SNP based lineage [51]. This lineage is distinct from the East Asian
tuberculosis lineages that includes Beijing type strains, which have been hypothesized to have increased virulence. Both SCG3b and SCG5 strains are expected to differ by a relatively small number of mutations because they are situated quite close to each other on SNP generated phylogenies, compared to more distantly related lineages such as the Beijing strains [26]. This suggests that only a small number of mutations may be responsible for the change in transmissibility observed in SC3b strains. These mutations may be discoverable in future whole genome sequencing studies.

One important limitation has been that most studies examining the virulence potential of clustered MTB isolates have used banked isolates with limited knowledge about the epidemiologic and clinical conditions present at the time the strains were cultured, and thus fail to consider potential confounders related to characteristics of the index case (e.g. cough severity, bacterial load, cavitations), environment (e.g. ventilation, crowding) or contact (e.g. age, BCG and proximity of contact) [10]. Household contact studies such as this one are a classic, well-accepted design to study MTB transmission dynamics in a well-characterized group of individuals with a defined infectious challenge, while controlling for host, environmental and bacterial confounders. However, contacts that are already infected at the time of initial ascertainment may have been infected by the source case or, in an intermediate prevalence country such as Brazil, infected in the community [52,53]. As observed in our study, this translates into age-dependent differences in prevalence of infection in contacts (i.e. higher frequency of TB exposures in adults as a result of wider social networks).

Our study has limitations. Whereas all patients were sputum smear- and culture-positive on solid media, we did not obtain quantitative culture data, which is known as a stronger surrogate of bacterial load and risk of infection. Between February 2008 and April 2009, we obtained a repeat TST in all contacts with TST ≥ 10 mm, including those living in IT households. As some IT households were excluded at the 1st TST after April 2009, some of which presumably would have been re-classified as

### Table 2. Distribution of households according to study *M. tuberculosis* transmission categories by study time point.

| Study time point | Final transmission category (2nd TST) | n (row %) |
|------------------|--------------------------------------|-----------|
|                  | Low                                  | Intermediate | High |
| n                | 41                                   | 23 (56)    | 11 (27)    | 7 (17)     |
| Intermediate     | 30                                   | NA         | 17 (57)    | 13 (43)    |
| High             | 53                                   | NA         | NA         | 53 (100)   |
| Totals           | 23                                   | 28          | 73         |

NA = Not applicable.  
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### Table 3. Household contacts with tuberculin skin test (TST) conversion from entry to week 8–12 by initial household transmission category and age of contact.

| TST conversion characteristic | Initial (1st TST) household transmission category | P* value |
|------------------------------|-----------------------------------------------|----------|
|                              | No. with TST conversion/No. at risk (%)       |          |
|                              | All                                           | Low      | Intermediate | High |
| Criterion 1³                | 50/271 (18.4)                                 | 36/189 (19.0) | 13/72 (18.1) | 1/10 (9.1) | 0.67 |
| ≤5 years                    | 5/43 (11.6)                                   | 4/32 (12.5) | 1/11 (9.1) | 0/0 (0.0) | 0.75 |
| 6–15 years                  | 23/86 (26.7)                                  | 17/63 (27.0) | 6/20 (30.0) | 0/3 (0.0) | 0.87 |
| ≥16 years                   | 22/142 (15.5)                                 | 15/94 (16.0) | 6/41 (14.6) | 1/7 (12.5) | 0.77 |
| Criterion 2²                | 50/221 (22.6)                                 | 36/159 (22.6) | 13/53 (24.5) | 1/9 (11.1) | 0.93 |
| ≤5 years                    | 5/42 (11.9)                                   | 4/31 (12.9) | 1/11 (9.1) | 0/0 (0.0) | 0.72 |
| 6–15 years                  | 23/76 (30.3)                                  | 18/57 (31.6) | 5/16 (31.3) | 0/3 (0.0) | 0.87 |
| ≥16 years                   | 22/103 (21.4)                                 | 14/61 (19.7) | 7/26 (26.9) | 1/6 (16.7) | 0.64 |

n/N are number of converters divided by the number at risk as defined by the various criteria below.

1 | TST conversion Criterion 1 (Brazilian guidelines): 1st TST < 10 mm; 2nd TST ≥ 10 mm; difference ≥ 10 mm. Does not include contacts with 1st TST ≥ 10 mm (407, 56%), and those with missing 1st (27, 3%) or 2nd TST (31, 4%) because they are not considered “at risk” by this criterion.

2 | TST conversion Criterion 2: 1st TST < 5 mm; 2nd TST ≥ 10 mm; difference ≥ 6 mm. Does not include contacts with 1st TST ≥ 5 mm (458, 65%), and those with missing 1st TST (2, 3%) or 2nd TST (31, 4%) because they are not considered “at risk” by this criterion.

*Score test from generalized estimating equation (GEE) estimation approach to logistic regression.

Only contacts “at risk” of TST conversion are included for each criterion.

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| Characteristic | Household Transmission Category | Unadjusted OR | P-value* | P-value* | Adjusted OR | P-value | P-value* |
|---------------|--------------------------------|---------------|----------|----------|-------------|---------|---------|
|               | n (column %)                  | [95% CI]      |          |          | [95% CI]    |         |         |
| Index cases   | Low                            |               |          |          |             |         |         |
|               | N=23                           |               |          |          |             |         |         |
|               | High                           |               |          |          |             |         |         |
| Age (per 10-year increment) | 23 | 73 | 0.90 | [0.62, 1.32] | 0.59 | 0.59 |
| Cough measurement at baseline | 21 | 70 | 1.36 | [1.13, 1.67] | 0.002 | 0.001 | 1.53 | [1.27, 1.86] | <0.001 |
| VACS (per unit increase) | 21 | 70 | 1.24 | [1.07, 1.45] | 0.007 | 0.005 |
| Chest radiograph |               |               |          |          |             |         |         |
| Normal/Minimal | 2 | 8.3 | 4 | [5.6] | 1 (ref) |           | 0.64E   |
| Moderate      | 13 | 56.5 | 35 | [49.3] | 1.34 | [0.11, 10.7E] | 1.00E   |
| Far advanced  | 8 | 34.8 | 32 | [45.1] | 2.00 | [0.25, 12.33E] | 0.78E   |
| Cavitations   |               |               |          |          |             |         |         |
| Absent        | 10 | 43.5 | 14 | [19.7] | 1 (ref) | 1 (ref) | 0.028 | 0.078 |
| Present       | 13 | 56.5 | 57 | [80.3] | 3.13 | [1.13, 8.69] | 0.027 | 3.50 | [0.97, 12.58] |
| Sputum AFB smear | 2+ | 6 | 26.1 | 12 | [16.4] | 1 (ref) |           | 0.31   |
|               | 3+ | 17 | 73.9 | 61 | [83.6] | 1.79 | [0.56, 5.38] | 0.30   |
| SNP cluster group | 3b | 6 | 27.3 | 9 | [12.7] | 1 (ref) | 1 (ref) | 0.21 | 0.40 |
| 5             | 15 | 68.2 | 55 | [77.5] | 2.44 | [0.72, 7.93] | 0.14 | 2.93 | [0.74, 11.64] | 0.13 |
| Other          | 1 | 4.5 | 7 | [9.9] | 5.33 | [0.70, 112.44] | 0.16 | 2.45 | [0.15, 39.91] | 0.53 |
| Household contacts | N=132 | | | | | | | |
| Age (years)   | <=5 | 21 | 15.9 | 48 | [11.5] | 1 (ref) | 1 (ref) | 0.47 | 0.17 |
|               | >5  | 37 | 28.0 | 127 | [30.5] | 1.50 | [0.78, 2.89] | 0.22 | 1.63 | [0.77, 3.46] | 0.23 |
| BCG scar      |               |               |          |          |             |         | 0.85   |
| Absent        | 21 | 15.9 | 76 | [18.3] | 1 (ref) |           |         |
| Present       | 107 | 81.1 | 326 | [78.4] | 0.84 | [0.43, 1.63] | 0.43 |
| Uncertain     | 4 | 3.0 | 14 | [3.4] | 0.97 | [0.23, 3.99] | 0.23 |
| Average exposure time to index case last 3 months (hours/day) | <=7 | 38 | 28.8 | 152 | [36.5] | 1 (ref) |           | 0.41   |
|               | >7  | 45 | 34.1 | 98 | [23.6] | 0.54 | [0.22, 1.33] | 0.18 |

Table 4. Unadjusted and Adjusted Odds Ratios of Index TB Case, Household Contact and Environmental Factors for Predicting High Mycobacterium tuberculosis Transmission Households.
| Characteristic | Household Transmission Category | Unadjusted OR | P-value* | Adjusted OR | P-value* |
|---------------|---------------------------------|---------------|----------|-------------|----------|
|               | n (column %)                    | [95% CI]      |          | [95% CI]    |          |
|               | Low                             |               |          |             |          |
| 13–18         | 31 (23.5)                       | 0.99 [0.34, 2.90] | 0.99     |             |          |
| ≥18           | 18 (13.6)                       | 0.60 [0.20, 1.79] | 0.36     |             |          |
| Environment   |                                 |               |          |             |          |
| Sleeping arrangement with index case | | | 0.51 |          |          |
| Same house    |                                 |               |          |             |          |
| Same room, same bed | 16 (12.1) | 70 (16.8) | 1 (ref) | 0.49 [0.18, 1.35] | 0.17 |
| Different room | 61 (46.2) | 178 (42.8) | 0.67 [0.32, 1.40] | 0.28 |
| Different house | 41 (31.1) | 138 (33.2) | 0.77 [0.28, 2.11] | 0.61 |

1
Household level data, except chest radiograph (see footnote E below), are modeled using logistic regression. Contact level data are modeled by using generalized estimating equations (GEE) approach to estimation for logistic regression with an independent working correlation matrix with standard errors obtained from a sandwich estimate of the variance.

2
Fit using a GEE approach to estimation for logistic regression with an independent working correlation matrix adjusted for all other covariates listed. Standard errors are from a sandwich estimator of the variance.

3
One participant whose SCG could not be distinguished between 5 or 6a was excluded.

*Wald (pairwise vs. reference) P-value.

+Global (omnibus) score P-value.

E = denotes exact Odds Ratios and P-values using an exact test of the parameters from exact logistic regression.

VACS = Visual analog cough scale; LCQ = Leicester cough questionnaire.

BCG = Bacille Calmette Guérin vaccine; TST = Tuberculin skin test; OR = Odds ratio; CI = Confidence Interval.

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HT, we cannot estimate the percentage of households that are HT within the study community. IT households are also underrepresented. However, because the initially HT group of households was large, the inclusion of additional households through conversion between the two TST applications into this group are unlikely to have altered the results of this study. Also, whereas our criteria to classify households were operationally driven, the definition of HT was robust in that 77% of HT households had 100% of households with TST ≥ 10 mm.

In conclusion, in this household contact study of culture confirmed, AFB + smear-positive TB patients in a Brazilian urban setting we observed variable MTB transmission within households. Our results provide strong evidence of differential transmission among otherwise clinically similar patients with advanced TB disease. Data presented here also suggest that these differences in infectiousness may be modulated by yet unknown interactions between the host and the infecting strain, resulting in weaker cough and possibly, less formation of cavities. Further studies will be needed to mechanistically advance these novel findings.

Supporting Information

Table S1 Additional characteristics of index tuberculosis cases, household contacts, and study dwellings at baseline by Mycobacterium tuberculosis transmission

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