Longitudinal genotyping surveillance of Mycobacterium tuberculosis in an area with high tuberculosis incidence shows high transmission rate of the modern Beijing subfamily in Japan

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Research paper

Longitudinal genotyping surveillance of *Mycobacterium tuberculosis* in an area with high tuberculosis incidence shows high transmission rate of the modern Beijing subfamily in Japan

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- Molecular epidemiology
- Beijing family
- VNTR

**ABSTRACT**

Tuberculosis (TB) is a severe and wide-spread infectious disease worldwide. The modern Beijing subfamily, one lineage of *M. tuberculosis*, reportedly has high pathogenicity and transmissibility. This study used a molecular epidemiological approach to investigate the transmissibility of the modern Beijing subfamily in the Airin area of Osaka City, Japan. During 2006–2016, we collected 596 *M. tuberculosis* clinical isolates in the Airin area, Osaka city, Japan. We analyzed the 24-locus variable number of tandem repeats typing optimized for the Beijing family of isolates, *M. tuberculosis* lineage, and patient epidemiological data. The proportion of the modern Beijing subfamily was significantly higher not only than previously obtained data for the Airin area: it was also higher than the nationwide in Japan. The rate of recent clusters, defined as a variable number of tandem repeats profile identified within two years, of the modern Beijing subfamily was significantly higher than that of the rate of recent clusters of the ancient Beijing subfamily. Results suggest that TB control measures formulated with attention to the modern Beijing subfamily might be an important benchmark to understanding recent TB transmission in the area.

1. Introduction

Tuberculosis (TB), an infectious bacterial disease caused by *Mycobacterium tuberculosis*, affects millions worldwide: 10.4 million people developed TB in 2016; 1.7 million died (WHO, 2017). Based on the World Health Organization End TB Strategy, which set a goal to end the TB epidemic by 2035 (WHO, 2014), every country must undertake strong countermeasures to end TB in accordance with their own country-specific TB situation. In Japan, 13.9 TB cases/100,000 population were reported in 2016. The incidence was 7.2–32.8 among prefectures and major cities. Osaka city, the third most populated city in Japan (SBJ; Ministry of Internal Affairs and Communications, 2017), reported the highest TB incidence. This high incidence is partly attributable to the Airin area, a residential area of many homeless people and day laborers in the city, which showed abnormally high TB incidence (e.g., est. 395 cases/100,000 population in 2016) (Matsumoto, 2015). Homeless people have been regarded as a high-risk group of TB (Bamrah et al., 2013; Hayward and Coker, 2000; Korzeniewska-Kosela et al., 2015). Countermeasures for high-risk groups are necessary to reduce TB incidence especially in urban areas, finally leading to an end to TB worldwide.

The Beijing family, within the East Asian lineage, is spreading worldwide from its origin in eastern Asia (Glynn et al., 2006; Parwati et al., 2010). The family is distinct from the other three major phylogenetic lineages of *M. tuberculosis* because of its genetic differences (Gagneux et al., 2006). Furthermore, the Beijing family had a
genetically evolved lineage, defined as the modern Beijing subfamily. This evolved sublineage could be classified by several genetic markers such as an insertion of the IS6110 sequence into namely the NTF region (Kremer et al., 2004; Mokrousov et al., 2005) and single nucleotide polymorphisms (Filliol et al., 2006; Nakanishi et al., 2013; Tsolaki et al., 2005). Earlier studies have demonstrated that the modern Beijing subfamily might have higher virulence and transmissibility than the ancient Beijing subfamily, not only by epidemiological studies but also by an experimental-based comparison (Hanekom et al., 2007a; Mokrousov et al., 2006; Ribeiro et al., 2014; van Laarhoven et al., 2013; Wada et al., 2009a). Considering prevention of the spread of TB, capturing the features of lineages and subfamilies is crucially important (Iwamoto et al., 2008, 2012; Langlois-Klassen et al., 2013; Luo et al., 2015; Maeda et al., 2014).

Reports of earlier studies have described that the modern Beijing subfamily is spreading globally (Bifani et al., 2002; Hanekom et al., 2007b; Iwamoto et al., 2012; Kato-Maeda et al., 2010; Mokrousov, 2013). In Japan, unlike the rest of the world, the ancient Beijing subfamily has been dominant (Iwamoto et al., 2008; Seto et al., 2015; Wada et al., 2009b), but the incidence of the modern Beijing subfamily is higher in younger age groups (Iwamoto et al., 2009; Seto et al., 2017). Therefore, distribution of the modern Beijing subfamily in Japan should be assessed adequately to call attention to the threat of TB expansion. For that purpose, elucidating the trend of the modern Beijing subfamily by dynamic tracking with long-term monitoring is fundamentally important.

Molecular epidemiology is an effective tool to discriminate isolates by genetic polymorphism and to verify infection links. Variable number of tandem repeats (VNTR) typing is a useful genotyping method to identify TB infections caused by the same isolate of M. tuberculosis (Supply et al., 2006). In Japan, > 75% of the clinical isolates belong to the Beijing family (Iwamoto et al., 2007; Murase et al., 2008; Wada et al., 2007; Yokoyama et al., 2007), 24Beijing-VNTR which was optimized for discriminating Beijing family isolates by 24 loci subset (Iwamoto et al., 2007, 2012). Transmission among TB patients can be surmised when clinical isolates show an indistinguishable VNTR profile (hereinafter cluster). In spite of the straightforward and cost-effective concept as a routine TB surveillance (de Beer et al., 2013), estimation of ongoing outbreaks based on VNTR clustering often tends to be overestimated because of its low discriminatory power. Misclustering of epidemiologically unrelated isolates occurs partly because of nature of M. tuberculosis, which has the potential to remain dormant in the human body after infection and which can cause reactivation of past latent TB infection.

Because of such a dormant phase of TB, cluster formation observed by genotyping of clinical isolates is not always correspondent to recent TB transmission (Lillebaek et al., 2002; Seto et al., 2017). To evaluate TB onset after recent transmission, we must address the feature of latent TB within two years of an initial infection. According to the paper, some tuberculin-positive persons at 10 years that 78% of people developed TB infection. Sutherland (1968) reported from a follow-up survey of TB cases that 40.0% received welfare support. Approximately 12,000 residents were day laborers or the homeless: many working poor people in all over Japan tend to migrate to this area seeking temporary employment. They live in cheap apartments, night shelters, and cheap hotels, but they often become homeless.

2. Materials and method

2.1. Study setting

The Airin area, occupying an area of 0.62 km² in the northeastern part of Nishinari-ward of Osaka city, is known as Japan’s largest urban poor area. In 2015, the area had 21,447 inhabitants, of whom 80.9% were male, 61.4% were elderly persons (≥60 years of age), and > 40.0% received welfare support. Approximately 12,000 residents were day laborers or the homeless: many working poor people in all over Japan tend to migrate to this area seeking temporary employment. They live in cheap apartments, night shelters, and cheap hotels, but they often become homeless.

2.2. Study population

We systematically collected M. tuberculosis clinical isolates from TB patients residing in the Airin area between January 2006 and December 2016. Demographic (sex, age) and clinical (site of disease, sputum smear status, and treatment history of TB) characteristics of the TB patients were connected with the isolates. We collected the isolates mainly from eight hospitals specializing in treatment of TB. This work was approved by the Ethics Committees of the Institute of Tropical Medicine, Nagasaki University (approval no. 17070172) and of Osaka Institute of Public Health (approval no. 1709-05).

2.3. VNTR analysis

M. tuberculosis isolates were suspended in H₂O and heated at 95 °C for 10 min. The supernatants of the heat-killed suspensions were used as PCR templates. 24Beijing-VNTR analysis was conducted according to reports of earlier studies (Iwamoto et al., 2007, 2012). The size of amplified PCR fragments was found using a capillary electrophoresis system (SV1210; Hitachi High-Technologies Corp., Tokyo, Japan) or a genetic analyzer (Applied Biosystems 3500 genetic analyzer; Thermo Fisher Scientific Inc., Waltham, MA, USA.) with the GeneMapper program (ver. 5.0) (Iwamoto et al., 2012; Wada et al., 2007). We calculated the number of repeats in each locus from the size of amplified PCR fragments (Iwamoto et al., 2007, 2012). All analyses were conducted at the Osaka Institute of Public Health.

2.4. Clustering analysis

A cluster was defined when two or more isolates showed indistinguishable 24Beijing-VNTR profiles. We used the definition as an indicator of the probability that transmission occurred within the Airin area during the surveillance. Generally, epidemiological links between clustered patients are necessary to prove actual transmission (Allix-Beguec et al., 2014; Seto et al., 2017). However, epidemiological information of patients in this area was difficult to obtain. Therefore, we introduced a definition of ‘recent clusters’ as an indicator of recent transmission. Because of this two-year time lag, only cases during 2008–2016 were used for the analysis of recent clusters. The rate of recent clusters in each year during 2008–2016 was calculated by dividing the total number of cases belonging to a recent cluster in one year by the number of cases for which VNTR analysis was conducted in that year (de Vries et al., 2007; Kong et al., 2002).

2.5. Estimation of M. tuberculosis lineage

We estimated the lineages of M. tuberculosis isolates using maximum a posteriori estimation with the 24Beijing-VNTR profile, as described in an earlier report (Seto et al., 2015). We integrated the estimation results of four sub-lineages (i.e., ST11/26, STK, ST3, and ST25/19) as the ancient Beijing subfamily. We used the results of the modern Beijing subfamily, the ancient Beijing subfamily, and the non-Beijing family in
this study. The population structure of \textit{M. tuberculosis} lineages in the Airin area was compared with those found in earlier studies of this area (Wada et al., 2009a) and others throughout Japan (Seto et al., 2015).

2.6. Statistical analysis

Kruskal–Wallis testing was applied to compare cluster sizes among \textit{M. tuberculosis} lineages. Adjusted residual analysis was used to assess the association between \textit{M. tuberculosis} lineages and patient age groups, and to compare the rates of cluster formation among \textit{M. tuberculosis} lineages. This analysis was also applied to compare the proportion of \textit{M. tuberculosis} lineages found in earlier studies with those found in this study. Cochran–Armitage tests were applied to investigate the trends of annual rates in cases belonging to recent clusters. To ascertain the association between recent cluster formation and epidemiologic features, we used logistic regression analysis to calculate the odds ratio and 95% CIs. We applied multivariate logistic regression analysis using age groups and \textit{M. tuberculosis} lineages as explanatory variables. We used backward stepwise for variable selection of multivariate analysis. We inferred results for which \( p < 0.05 \) as statistically significant. All statistical analyses were conducted using software (R ver. 3.4.3; R Foundation for Statistical Computing, Vienna, Austria).

3. Results

3.1. Characteristics of the study population

During 2006–2016, 1325 cases were registered in the Airin area, Osaka City, Japan; 810 of the cases were culture positive. We investigated 596 isolates (45.0%) of the 1325 cases (73.6% of 810 culture positive cases) for this study.

The 596 patients had mean ± SD age of 63.3 ± 11.1 (range 22–92) years. Most patients were male (579 patients, 97.1%), Japanese (594 patients, 99.7%) and undergoing primary TB treatment (527 patients, 88.4%). Pulmonary TB (591 patients, 99.2%) and sputum smear-positive (480 patients, 80.5%) also had high proportions. Multidrug-resistant (MDR) patients were four (0.67%). The proportion of recent cluster formation was significantly higher by Adjusted residual analysis (\( p < 0.05 \)).

3.2. Distribution of \textit{M. tuberculosis} lineages

All of the 596 isolates were classified into three phylogenetic groups: the non-Beijing family (117 (19.6%) isolates), the ancient Beijing subfamily (251 (42.1%) isolates), and the modern Beijing subfamily (228 (38.3%) isolates) (Table 1). It is noteworthy that no statistical significance by age group was found in the proportion of the modern Beijing subfamily.

3.3. Overall TB transmission

Genotyping analysis based on \( 24_{\text{Beijing}} \)-VNTR typing revealed that 241 (40.4%) of the 596 isolates formed 56 clusters during 2006–2016 (Supplementary Table S1, Fig. 1). We compared the proportion of cluster formation in cases of non-Beijing family (nBJ), the ancient Beijing subfamily (Ancient), and the modern Beijing subfamily (Modern): white circles represent clusters; black bars show median cluster size in each lineage.
using TB patient characteristics (Table 3). Patients 49 years old or younger had a significantly high odds ratio than patients 70 years old or older only in univariate analysis. In both univariate and multivariate analyses, sputum smear-positive cases showed a significantly higher odds ratio than sputum smear-negative cases. Also, the ancient Beijing subfamily had a significantly lower odds ratio than the modern Beijing subfamily, but statistical significance was not found between the modern Beijing subfamily and the non-Beijing family.

Finally, risk factors for recent TB transmission were investigated using TB patient characteristics (Table 3). Patients 49 years old or younger had a significantly high odds ratio than patients 70 years old or older only in univariate analysis. In both univariate and multivariate analyses, sputum smear-positive cases showed a significantly higher odds ratio than sputum smear-negative cases. Also, the ancient Beijing subfamily had a significantly lower odds ratio than the modern Beijing subfamily, but statistical significance was not found between the modern Beijing subfamily and the non-Beijing family.

4. Discussion

In this study, we monitored transmission activities of the modern Beijing subfamily in the most prevalent TB area of Japan. We verified the trend of the modern Beijing subfamily along with genotyping approach and patients' epidemiological data, during more than a decade. In spite of recent reports describing the modern Beijing subfamily originating in northern China (Luo et al., 2015) has become dominant worldwide (Bifani et al., 2002; Hanekom et al., 2007b; Ivamoto et al., 2012; Kato-Maeda et al., 2010; Mokrousov, 2013; Yin et al., 2016), the transmissibility of the modern Beijing isolates remains unclear. In a basic research, the modern Beijing subfamily's survival strategy was implied that induced a lower level of proinflammatory cytokines compared with the ancient Beijing subfamily, could escape from the host immune system (van Laarhoven et al., 2013). Our findings indicate that the modern Beijing subfamily deserves attention as a threat of transmission in a high-TB incidence area.

Regarding the overall trend of TB in the Airin area, the rate of recent clusters has been decreasing (Fig. 2). Furthermore, VNTR clusters were less detected during the survey period in the latter half (21.4%) (Supplementary Table S1). Those results indicate that recent isolates tended not to cause direct transmission in this area. The reduction of TB transmission may have been attributable to vigorous implementation of TB control for the area. Since 2001, the Osaka City Public Health Center has taken measures such as (1) patient supports daily Directly Observed Treatment: health staff observe TB patients to swallow TB medicine in front of them to make sure they are taking medicine daily, and (2) TB screening by chest x-ray examination for residents to find new TB cases efficiently. The activities have been strengthened since 2012 to assure to provide free medical services and residences/rooms/beds for all TB patients and TB suspects in this area. As results, from 2013 to 2016, lost of follow-up rate reduced from 4.5% to 2.8%, and proportion of patients detected by TB screening increased from 17.7% to 24.7% (data not shown). Furthermore, MDR rate of smear-positive patients reduced from 2.53% (35/1383) in 2001–2008 (Shimouchi et al., 2013) to 0.47% (1/214) in 2013–2016 (data not shown). However in order to cope with transmission, early case finding of TB patients among the homeless and contact examination among residents of apartments should be further strengthened and expanded. Efforts to control TB in high incidence areas is indispensable to achieve the global eradication of TB (WHO, 2015). Molecular epidemiological monitoring is expected to become more important as the incidence rate decreases.

The modern Beijing subfamily was estimated as exhibiting high transmissibility in the Airin area from the viewpoint that the cluster size was regarded as indicating high propagation of isolates (Ypma et al., 2013). The largest cluster was composed of non-Beijing, but three of the five large clusters belonged to the modern Beijing subfamily.

### Table 3

Risk factors for the recent transmission of *M. tuberculosis* in the Airin area, Osaka City, Japan, 2008—2016.

<table>
<thead>
<tr>
<th>Patient characteristic</th>
<th>24<em>Beijing</em>-VNTR profile, no. (%)</th>
<th>Odds ratio (95% CI)*&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Univariate</th>
<th>Multivariate&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age group</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤49</td>
<td>36 (10.2)</td>
<td>19 (15.1)</td>
<td>2.04 (1.03–4.05)</td>
<td>-</td>
</tr>
<tr>
<td>50–59</td>
<td>71 (20.1)</td>
<td>31 (24.6)</td>
<td>1.69 (0.94–3.02)</td>
<td>-</td>
</tr>
<tr>
<td>60–69</td>
<td>130 (36.8)</td>
<td>46 (36.5)</td>
<td>1.37 (0.81–2.31)</td>
<td>-</td>
</tr>
<tr>
<td>≥70</td>
<td>116 (32.9)</td>
<td>30 (23.8)</td>
<td>1.0</td>
<td>-</td>
</tr>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>9 (2.5)</td>
<td>6 (4.8)</td>
<td>1.91 (0.67–5.48)</td>
<td>-</td>
</tr>
<tr>
<td>Male</td>
<td>344 (97.5)</td>
<td>120 (95.2)</td>
<td>1.0</td>
<td>-</td>
</tr>
<tr>
<td><strong>Disease site</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pulmonary, sputum smear-positive</td>
<td>276 (78.2)</td>
<td>110 (87.3)</td>
<td>2.14 (1.16–3.94)</td>
<td>2.21 (1.18–4.12)</td>
</tr>
<tr>
<td>Pulmonary, sputum smear-negative</td>
<td>75 (21.2)</td>
<td>14 (11.1)</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Extrapulmonary</td>
<td>2 (0.6)</td>
<td>2 (1.6)</td>
<td>5.36 (0.70–41.25)</td>
<td>4.11 (0.52–32.51)</td>
</tr>
<tr>
<td><strong>Treatment history</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary</td>
<td>311 (88.1)</td>
<td>116 (92.1)</td>
<td>1.57 (0.76–3.22)</td>
<td>1.71 (0.81–3.61)</td>
</tr>
<tr>
<td>Retreatment</td>
<td>42 (11.9)</td>
<td>10 (7.9)</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td><strong>M. tuberculosis lineage</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Beijing</td>
<td>66 (18.7)</td>
<td>24 (19.0)</td>
<td>0.62 (0.36–1.08)</td>
<td>0.62 (0.36–1.10)</td>
</tr>
<tr>
<td>Ancient Beijing</td>
<td>171 (48.4)</td>
<td>34 (27.0)</td>
<td>0.34 (0.21–0.55)</td>
<td>0.34 (0.21–0.55)</td>
</tr>
<tr>
<td>Modern Beijing</td>
<td>116 (32.9)</td>
<td>68 (54.0)</td>
<td>1.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>

* CI’s that do not overlap the null value of odds ratio = 1 are shown in bold.

<sup>a</sup> Adjusted for the other factors used in the multivariate model.

<sup>b</sup> No variables.
Furthermore, although no significant difference was observed, the median cluster size of the modern Beijing subfamily was larger than that of the other lineages (Fig. 1). One aspect of the social background in the Airin area is that low-income people who have difficulty finding jobs in their hometown immigrate from throughout Japan. Further efforts to incorporate patient information into research design must be undertaken to resolve whether increase of the modern Beijing subfamily is related to the inflow of people or stable circulation of the subfamily within the area.

Expansion of the modern Beijing subfamily was not limited to certain age groups in our setting (Table 1). In earlier studies, the modern Beijing subfamily was found to be significantly predominant in younger age groups in Japan [Iwamoto et al., 2009; Seto et al., 2017]. In contrast to that tendency, the subfamily was isolated from TB patients in the Airin area irrespective of age groups in this study (Table 1). The result might be related to a common behavior of persons living in this area; they usually do not belong to a specific community and spend long periods in close and overcrowded spaces (e.g., shelters, libraries, and resting places) irrespective of their age.

Our results suggest that the modern Beijing subfamily affects recent TB transmission in the Airin area. Multivariate analysis for recent cluster formation showed the modern Beijing subfamily cause more recent transmission than the ancient Beijing subfamily (Table 3). Although the same tendency was not confirmed between the non-Beijing family and the modern Beijing subfamily, the largest cluster of the non-Beijing family (Supplementary Table S1) might influence the non-significant result. Except for that case, the modern Beijing subfamily was constantly transmitted within the area. This assumption would also be suggested by the proportion of the modern Beijing subfamily in the Airin area was clearly increased over previous studies [Wada et al., 2009a,b].

<table>
<thead>
<tr>
<th>Area</th>
<th>Study period</th>
<th>n (%)</th>
<th>No. of isolates (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Non-Beijing</td>
</tr>
<tr>
<td>Previous study of Airin area</td>
<td>2002–2004</td>
<td>274 (100)</td>
<td>61 (22.3)</td>
</tr>
<tr>
<td>All over Japan</td>
<td>2004–2013</td>
<td>1245 (100)</td>
<td>310 (24.9)</td>
</tr>
<tr>
<td>This study</td>
<td>2006–2016</td>
<td>596 (100)</td>
<td>117 (19.6)</td>
</tr>
</tbody>
</table>

* Wada et al. (2009a,b).
* Seto and Wada et al. (2015).
* p < 0.05, Adjusted residual analysis between This study or Previous study of Airin area.
* p < 0.05, Adjusted residual analysis between This study or All over Japan.
* p < 0.05, Adjusted residual analysis between This study or All over Japan.

5. Conclusion

A longitudinal molecular epidemiologic monitoring based on VNTR clusters within two years revealed that transmission of TB was decreasing in the Airin area. However, TB incidence in this area was still high (395 cases/100,000 population in 2016) and active transmission still occurred continuously. Under such circumstances, the modern Beijing subfamily was confirmed as a risk factor of recent TB transmissions compared with other lineages in the Airin area. Furthermore, the proportion of this subfamily was found to be increasing in this area. These results suggest that the modern Beijing subfamily represents an epidemic threat in areas with high TB incidence. It might require careful consideration of further strengthening and expansion for TB control measures.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.meegid.2018.09.014.

References


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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.meegid.2018.09.014.


