Directly Observed Therapy and Improved Tuberculosis Treatment Outcomes in Thailand

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Abstract

Background: The World Health Organization (WHO) recommends that tuberculosis (TB) patients receive directly observed therapy (DOT). Randomized controlled trials have not consistently shown that this practice improves TB treatment success rates. In Thailand, one of 22 WHO-designated high burden TB countries, patients may have TB treatment observed by a health care worker (HCW), family member, or no one. We studied whether DOT improved TB treatment outcomes in a prospective, observational cohort.

Methods and Findings: We prospectively collected epidemiologic data about TB patients treated at public and private facilities in four provinces in Thailand and the national infectious diseases hospital from 2004–2006. Public health staff recorded the type of observed therapy that patients received during the first two months of TB treatment. We limited our analysis to pulmonary TB patients never previously treated for TB and not known to have multidrug-resistant TB. We analyzed the proportion of patients still on treatment at the end of two months and with treatment success at the end of treatment according to DOT type. We used propensity score analysis to control for factors associated with DOT and treatment outcome. Of 8,031 patients eligible for analysis, 24% received HCW DOT, 59% family DOT, and 18% self-administered therapy (SAT). Smear-positive TB was diagnosed in 63%, and 21% were HIV-infected. Of patients either on treatment or that defaulted at two months, 1601/1636 (98%) patients that received HCW DOT remained on treatment at two months compared with 1096/1268 (86%) patients that received SAT (adjusted OR [aOR] 3.8; 95% confidence interval [CI] 2.4–6.0) and 3782/3987 (95%) patients that received family DOT (aOR 2.1; CI 1.4–3.1). Of patients that had treatment success or that defaulted at the end of treatment, 1369/1477 (93%) patients that received HCW DOT completed treatment compared with 744/1074 (69%) patients that received SAT (aOR 3.3; CI 2.4–4.5) and 3130/3529 (89%) patients that received family DOT (aOR 1.5; 1.2–1.9). The benefit of HCW DOT compared with SAT was similar, but smaller, when comparing patients with treatment success to those with death, default, or failure.

Conclusions: In Thailand, two months of DOT was associated with lower odds of default during treatment. The magnitude of benefit was greater for DOT provided by a HCW compared with a family member. Thailand should consider increasing its use of HCW DOT during TB treatment.

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Introduction

Despite the widespread availability of cheap, effective treatment, tuberculosis (TB) remains a major cause of severe illness and death, with an estimated nine million new cases and two million deaths occurring annually.[1] One barrier to global TB control is the long duration of TB treatment—a minimum of six months—which frequently results in patients taking their medications erratically or not at all.[2] Non-adherence to TB medications decreases the chances of cure, increases the risk of relapse after treatment, and selects for drug-resistant TB strains.[3] Directly observing TB patients taking their anti-TB therapy, either daily or several times per week, was first piloted in the 1950s as a way to insure adherence and treatment completion.[2] In 1994, based on the reported success of directly observed therapy (DOT) in increasing treatment completion rates and preventing drug resistance, the World Health Organization (WHO) adopted DOT as a principal component of its global TB control
strategy.[4] Current technical manuals define DOT as direct supervision of “medication ingestion…by a treatment supporter who is acceptable and accountable to the patient and to the health system.”[5]

Although WHO and other international agencies strongly advocate DOT, controversy remains whether its benefits have been proven. Randomized controlled trials (RCT) have shown either modest or no benefit of DOT in improving TB treatment success rates, and a meta-analysis of 10 RCTs concluded that the evidence base for WHO’s DOT policy is insufficient.[6] Advocates of DOT have argued that the RCTs and this meta-analysis evaluated the wrong endpoint. The effectiveness of DOT, it has been argued, should be judged by how well it prevents drug resistance, specifically to rifampin, and not by improvements in treatment success rates.[7] Others contend that the scientific literature supporting DOT has been weakened by studies involving “sloppy” DOTs; the RCTs, it has been argued, were under-powered to show an improvement in treatment success rates, because the programs studied had sub-standard TB programs.[8]

While RCTs are considered the gold standard for measuring the efficacy of a biomedical intervention, prospective observational studies are required to evaluate the effectiveness of an intervention applied to a large population in uncontrolled (i.e., real world) settings.[9] Such studies are particularly necessary for an intervention, such as DOT for TB treatment, that involves multiple components of the health system and relies predominantly on government health facilities in poor countries. Thailand is a low middle-income country with the 17th largest burden of TB in the world.[1] Despite official adoption of the WHO TB control strategy in 1997, TB rates in Thailand have failed to decline, likely due to a generalized HIV epidemic and sub-optimal treatment success rates.[10] In Thailand, patients are treated in both the public and private sector, and different strategies for DOT, including no DOT, are implemented. Using data prospectively collected over two years, we evaluated the impact of different DOT strategies on treatment outcomes in a large, diverse cohort of TB patients.

Methods

Data Collection
In 2003, the United States Centers for Disease Control and Prevention (U.S. CDC) began collaborating with the Thailand Ministry of Public Health (MOPH), Japan’s Research Institute for Tuberculosis (RIT), four provinces in Thailand (Bangkok, Ubon-Ratchathani, Phuket, Chiang Rai), and the national infectious diseases hospital (in Nonthaburi province) on the Thailand TB Active Surveillance Network, a demonstration project involving enhanced surveillance, monitoring, evaluation, and treatment of TB in Thailand.[11]

For all patients with a diagnosis of TB in the national infectious diseases hospital or any public or private facility in the four provinces, public health staff recorded standardized epidemiologic data, collected sputum specimens for microbiologic testing, and offered HIV counseling and testing. Patient data was collected prospectively from routine medical and laboratory records and entered into an electronic database. Patient outcomes were recorded through the end of TB treatment, which was usually about six months after registration.

Patient Population
All persons registered for TB treatment were considered TB patients, consistent with WHO guidelines.[12] In this study, patients were eligible for analysis if they were registered for TB treatment between 1 October 2004 to 30 September 2006, were diagnosed with pulmonary TB, were not previously treated for TB or transferred in from a different TB program, were not known to have multidrug-resistant TB (MDR-TB), and had data recorded about their treatment observer. We classified patients with extra-pulmonary TB as ineligible, because the duration of treatment, drug regimen, and classification of outcomes, such as failure, vary depending on the location of disease.[12] We classified patients with previous TB treatment or known MDR-TB as ineligible, because such patients are known to have substantially different treatment outcomes than patients never previously treated.[13] Eligible patients were excluded from the analysis of treatment outcomes if their TB diagnosis was changed after registration, they were missing data about treatment status at two months (for the two month outcome analysis), or they were missing data about their final treatment outcome (for the end of treatment analysis).

For this study, patients with an outcome of “transferred out” or patients still on treatment at the time of this analysis were considered to have missing outcome data.

Definitions
We used standard WHO definitions to categorize patients according to previous TB treatment history, type of TB, and treatment outcome, and we classified any death which occurred during TB treatment as a TB death.[12] Consistent with WHO recommendations, sputum culture was not used to evaluate treatment outcome.[12]

In the database, public health staff recorded the type of treatment observer used during the first two months of TB treatment. Staff were instructed to classify patients as having health care worker (HCW) DOT if the patient had ingestion of anti-TB medicine observed by a HCW at least five times per week, and they were instructed to classify patients as having family DOT if the patient had a family member educated about TB treatment who was responsible for observing and recording ingestion of anti-TB medicine. Data was only recorded about the type of observer used during the first two months of treatment, because Thai national guidelines only require DOT during the period in which four drugs are administered. In actual practice, some facilities that provided DOT did so throughout treatment. The decision to allocate patients to different forms of DOT was made by individual health care providers; no data was collected about why different strategies were used in different patients.

Data Analysis
We divided patients into three groups: HCW DOT, family DOT, or self-administered treatment (SAT). We compared the association between type of DOT received and treatment outcome at two months and at the end of TB treatment. For the two month outcome analysis, patient outcomes were divided into on treatment, died, or defaulted. We compared the proportion of patients still on treatment at two months versus those that defaulted and the proportion still on treatment versus those that either died or defaulted, according to DOT type. For the end of treatment outcome analysis, patient outcomes included successful treatment (defined as cured or completed treatment), died, defaulted, or failed. We compared the proportion of patients successfully treated versus those that died, defaulted, or failed and treatment and the proportion of patients successfully treated versus those that defaulted, according to DOT type. In both the two month and end of treatment analysis, default was analyzed separately, because DOT is postulated to help reduce rates of default.[14]
In bivariate analysis, we calculated the odds ratio (OR) and 95% confidence interval (CI) for factors associated with the use of HCW, family DOT, or SAT. Statistical significance was defined as p<0.05. Because some groups of patients were more likely to receive a specific type of DOT and some factors associated with DOT use were also associated with treatment outcomes, we analyzed the association between DOT and treatment outcomes using propensity score analysis.

Propensity score analysis is used when the baseline characteristics of patients in two exposure groups (for example, those receiving HCW DOT vs. those receiving family DOT) are very different.[15] In observational studies, propensity score analyses can produce a more accurate estimate of the true association between an intervention (e.g., DOT) and an outcome (e.g., treatment success) by combining factors associated with the intervention into a composite variable, known as the propensity score, and by dividing the study population into strata that differ with respect to the likelihood of receiving the intervention, but are mostly equal with respect to other covariates.[15,16] In this study, we first developed a multivariate logistic regression model of factors associated with DOT, constructed propensity scores based on these factors, and then divided the patient population into equally sized quintiles based on their propensity to receive DOT. We used logistic regression to calculate adjusted odds ratios for the association between treatment outcomes and DOT type controlling for the DOT propensity quintile. We used an identical approach for 12 different analyses, i.e., three exposure comparisons (HCW vs. family DOT; family DOT vs. SAT; and HCW vs. family DOT) analyzed in four different patient subsets (success vs. death, default, or failure at the end of TB treatment; success vs. default at the end of TB treatment; on treatment vs. death or default at two months; and on treatment vs. default at two months).

Because there were variables consistently associated with successful treatment or type of DOT in all 12 analysis, we included these in the calculation of propensity scores: age, gender, marital status, Thai nationality, mobility (defined as not residing for at least three of last six months in the same district), living in an urban district, chronic cough, history of injection drug use, history of being in prison, history of previous isoniazid (INH) preventive therapy, diabetes, HIV infection, having a cavity on chest radiograph, sputum culture positive for *Mycobacterium tuberculosis*, treatment with a standardized regimen (e.g., WHO Category I), and quarter of enrollment.

### Ethical Review

The protocol for this demonstration project underwent ethical review by the Thailand Ministry of Public Health and CDC and was found to be surveillance and public health program implementation, not human subjects research requiring oversight by an institutional review board.

### Results

#### Patients analyzed

Of 14,354 patients recorded in surveillance, 8,031 (56%) were eligible for the analysis.[Figure 1] The most common reason for non-eligibility was extra-pulmonary TB (22%). The end-of-
Table 1. Characteristics of pulmonary TB patients eligible for analysis, stratified by type of observer during first two months of TB treatment.

| Characteristics                        | Treatment observer type | Total          |
|----------------------------------------|-------------------------|----------------|
|                                        | Health care worker | Family | Self-Administered | (N = 8,031) |
|                                        | (N = 1,900) | (N = 4,725) | (N = 1,406) |
| Type of pulmonary TB                   |                        |                |                |
| Smear-positive                         | 1230 (65) | 3072 (65) | 730 (52) | 5032 (63) |
| Smear-negative                         | 662 (35) | 1622 (34) | 457 (32) | 2741 (34) |
| Smear-unknown                          | 8 (0) | 31 (1) | 219 (16) | 258 (3) |
| Age                                    |                        |                |                |
| 0–14                                   | 20 (1) | 99 (2) | 10 (1) | 129 (2) |
| 15–44                                  | 1180 (62) | 2077 (44) | 948 (67) | 4205 (52) |
| 45–64                                  | 503 (27) | 1435 (30) | 317 (23) | 2255 (28) |
| >65                                    | 197 (10) | 1110 (24) | 127 (9) | 1434 (18) |
| Missing                                 | 0 (0) | 4 (0) | 4 (0) | 8 (0) |
| Gender                                 |                        |                |                |
| Male                                   | 1330 (70) | 3091 (65) | 880 (63) | 5301 (66) |
| Female                                 | 570 (30) | 1632 (35) | 526 (37) | 2728 (34) |
| Missing                                 | 0 (0) | 2 (0) | 0 (0) | 2 (0) |
| Nationality                           |                        |                |                |
| Thai                                   | 1747 (92) | 4449 (94) | 1162 (83) | 7358 (92) |
| Non-Thai                               | 153 (8) | 269 (6) | 241 (17) | 663 (8) |
| Missing                                 | 0 (0) | 7 (0) | 3 (0) | 10 (0) |
| Marital status                         |                        |                |                |
| Married                                | 1091 (57) | 3068 (65) | 543 (39) | 4702 (59) |
| Non-Married                            | 798 (42) | 1610 (34) | 632 (45) | 3040 (38) |
| Missing                                 | 11 (1) | 47 (1) | 231 (16) | 289 (3) |
| Living in an urban district            |                        |                |                |
| Urban district                         | 877 (46) | 1608 (34) | 823 (59) | 3308 (41) |
| Non-urban district                     | 1000 (53) | 3001 (64) | 483 (34) | 4484 (56) |
| Missing                                 | 23 (1) | 116 (2) | 100 (7) | 239 (3) |
| Mobile population*                     |                        |                |                |
| Non-mobile                             | 1560 (82) | 3738 (79) | 467 (33) | 5765 (72) |
| Mobile                                 | 315 (17) | 836 (18) | 798 (57) | 1949 (24) |
| Missing                                 | 25 (1) | 151 (3) | 141 (10) | 317 (4) |
| Ever treated with isoniazid preventive therapy (IPT) | | | |
| Previously treated with IPT            | 22 (1) | 28 (1) | 7 (1) | 57 (1) |
| Not ever treated with IPT              | 1876 (99) | 4688 (99) | 1354 (96) | 7918 (98) |
| Missing                                 | 2 (0) | 9 (0) | 45 (3) | 56 (1) |
| Cough lasting >2 weeks at time of diagnosis | | | |
| Cough >2 weeks                         | 1404 (74) | 3258 (69) | 805 (57) | 5467 (68) |
| No cough >2 weeks                      | 487 (26) | 1389 (29) | 317 (23) | 2193 (27) |
| Missing                                 | 9 (0) | 78 (2) | 284 (20) | 371 (5) |
| Ever used injection drugs              |                        |                |                |
| Ever used injection drugs              | 80 (4) | 88 (2) | 36 (3) | 204 (3) |
| Not ever used injection drugs          | 1769 (93) | 4545 (96) | 779 (55) | 7093 (88) |
| Missing                                 | 51 (3) | 92 (2) | 591 (42) | 734 (9) |
| In jail or prison                      |                        |                |                |
| Previously in jail or prison           | 80 (4) | 41 (1) | 7 (0) | 128 (2) |
| Not in jail/prison                     | 1805 (95) | 4671 (99) | 1006 (72) | 7482 (93) |
| Missing                                 | 15 (0) | 13 (0) | 393 (28) | 421 (5) |
| Characteristics                          | Treatment observer type | Total |
|-----------------------------------------|-------------------------|-------|
|                                         | Health care worker      | Family| Self-Administered| No. (%) |
|                                         | (N = 1,900)             | (N = 4,725) | (N = 1,406) |
| Living in migrant or refugee camp       |                         |       |
| In camp                                 | 83 (4)                  | 19 (0)| 10 (1)          | 112 (1) |
| Not in camp                             | 1765 (93)               | 4626 (98)| 844 (60) | 7235 (90) |
| Unknown                                 | 52 (3)                  | 80 (2)| 552 (39)        | 684 (9) |
| Facility that made diagnosis            |                         |       |
| Private health facility                 | 46 (3)                  | 229 (5)| 608 (43)        | 883 (11) |
| Government health facility              | 1853 (97)               | 4494 (95)| 797 (57) | 7144 (89) |
| Missing                                 | 1 (0)                   | 2 (0)| 1 (0)           | 4 (0) |
| Facility that provide treatment         |                         |       |
| Private health facility                 | 10 (1)                  | 172 (4)| 604 (43)        | 786 (10) |
| Government health facility              | 1889 (9)                | 4551 (96)| 802 (57) | 7242 (90) |
| Missing                                 | 1 (0)                   | 2 (0)| 0 (0)           | 3 (0) |
| Diabetes mellitus                       |                         |       |
| Diabetes                               | 63 (3)                  | 236 (5)| 37 (3)          | 336 (4) |
| No diabetes                            | 1459 (77)               | 3881 (82)| 886 (63) | 6226 (78) |
| Missing                                | 378 (20)                | 608 (13)| 483 (34)        | 1469 (18) |
| HIV status                             |                         |       |
| Positive                               | 433 (23)                | 983 (21)| 300 (21)        | 1716 (21) |
| Negative                               | 1322 (70)               | 2834 (60)| 365 (26) | 4521 (56) |
| Missing                                | 145 (7)                 | 908 (19)| 741 (53)        | 1794 (22) |
| Chest radiograph                       |                         |       |
| Normal                                 | 15 (1)                  | 76 (1)| 34 (2)          | 125 (2) |
| Not performed or results missing       | 103 (5)                 | 550 (12)| 63 (5)          | 716 (9) |
| Abnormal                               | 1782 (94)               | 4099 (87)| 1309 (93) | 7190 (89) |
| Presence of a cavity                   | 490 (28)                | 1164 (28)| 272 (21)        | 1926 (22) |
| Sputum culture result                  |                         |       |
| Growth of MTB                          | 1118 (59)               | 2297 (49)| 349 (25)        | 3764 (47) |
| No growth                              | 400 (21)                | 686 (14)| 127 (9)         | 1213 (15) |
| Not performed, contaminated, or grew NTM | 382 (20)             | 1742 (37)| 930 (66)        | 3054 (38) |
| Initial treatment prescribed           |                         |       |
| CAT I (2HRZE/4HR)                      | 1785 (94)               | 4454 (94)| 1126 (80) | 7365 (92) |
| Other regimens                         | 115 (6)                 | 271 (6)| 280 (20)        | 666 (8) |
| Period of the year                     |                         |       |
| Oct.–Dec.                              | 492 (26)                | 995 (21)| 289 (20)        | 1776 (22) |
| Jan.–Mar.                              | 506 (26)                | 1211 (26)| 419 (30)        | 2136 (27) |
| Apr.–Jun.                              | 454 (24)                | 1331 (28)| 376 (27)        | 2161 (27) |
| Jul.–Sep.                              | 447 (24)                | 1188 (25)| 322 (23)        | 1957 (24) |
| Missing                                | 1 (0)                   | 0 (0)| 0 (0)           | 1 (0) |
| Treatment outcome at the end of intensive phase |             |       |
| Smear-negative                         | 1295 (68)               | 2937 (62)| 416 (30)        | 4648 (58) |
| Smear-positive                         | 51 (3)                  | 131 (3)| 46 (3)          | 228 (3) |
| Died                                   | 148 (8)                 | 412 (9)| 48 (3)          | 608 (8) |
| Default                                | 35 (2)                  | 206 (4)| 173 (12)        | 414 (5) |
| Transferred out                        | 86 (4)                  | 177 (4)| 70 (5)          | 333 (4) |
| Change of diagnosis                    | 29 (2)                  | 118 (3)| 10 (1)          | 157 (2) |
| On treatment, smear unknown®          | 256 (3)                 | 721 (15)| 640 (46)        | 1617 (20) |
| Missing                                | 0 (0)                   | 23 (0)| 3 (0)           | 26 (0) |
treatment analysis included 7,070 patients, because we excluded 961 (12%) eligible patients that were recorded as still being on treatment, as transferred out, or as changed diagnosis. The two month outcome analysis included 7,515 patients, because we excluded 516 (6%) eligible patients that had an outcome recorded.

Table 1. cont.

| Characteristics          | Treatment observer type | Total     |
|--------------------------|-------------------------|-----------|
|                          | Health care worker (N = 1,900) | Family (N = 4,725) | Self-Administered (N = 1,406) |
|                          | (%)                      | (%)       | (%)       |
| Cured                    | 877 (46)                | 1937 (41) | 205 (14)  |
| Completed                | 492 (26)                | 1198 (25) | 544 (39)  |
| Failure                  | 17 (1)                  | 74 (2)    | 13 (1)    |
| Died                     | 222 (12)                | 583 (12)  | 67 (5)    |
| Default                  | 108 (6)                 | 401 (8)   | 332 (23)  |
| Transfer out             | 121 (6)                 | 245 (5)   | 100 (7)   |
| Change of diagnosis      | 29 (1)                  | 117 (2)   | 11 (1)    |
| On treatment             | 34 (2)                  | 170 (4)   | 134 (10)  |

*Mobile was defined as not living in the same district for at least three of the past six months.

MTB denotes Mycobacterium tuberculosis, and NTM denotes non-tuberculous mycobacteria.

Patients who were on treatment, but had missing data about whether their sputum smears were positive or negative.

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Outcomes at end of two months

When limited to patients that were on treatment, died, or defaulted within two months of starting TB treatment, 1,601 (90%) of 1,784 patients that received HCW DOT were still on treatment at two months, compared with 1,996 (89%) of 3,167 patients that received SAT (OR 1.1; CI, 0.9–1.4), when adjusted for propensity score, the magnitude of association decreased, and was not statistically significant (adjusted OR [aOR] 1.2; CI, 0.9–1.6).

When we restricted the analysis to evaluate the impact of DOT on default at two months, HCW DOT was strongly associated with being on treatment at two months compared with SAT or family DOT, because patients who received HCW DOT had a faster rate of treatment success or default at the end of TB treatment, compared with patients that received SAT or family DOT.

When we restricted the analysis to evaluate the impact of DOT on death at two months, HCW DOT was strongly associated with being on treatment at two months compared with SAT or family DOT.

Outcomes at end of treatment

At the end of TB treatment, treatment success was associated with SAT (aOR 1.6; CI, 1.3–2.0) or family DOT (aOR 1.3; CI, 1.1–1.5) compared with patients that received SAT.

When we restricted the analysis to evaluate the impact of DOT on only death at the end of TB treatment, HCW DOT was strongly associated with treatment success compared with patients that received SAT.

When we restricted the analysis to evaluate the impact of DOT on only default at the end of TB treatment, HCW DOT was strongly associated with treatment success compared with patients that received SAT.

When we restricted the analysis to evaluate the impact of DOT on only death at the end of TB treatment, HCW DOT was strongly associated with treatment success compared with patients that received SAT.
Table 2. Characteristics of eligible TB patients, stratified by inclusion or exclusion from two month and end of treatment analysis.

|                          | At Two Months |                           | At End of Treatment |                           |
|--------------------------|---------------|---------------------------|---------------------|---------------------------|
|                          | Excluded      | Included                  | Excluded            | Included                  |
|                          | No. (%) (N=516)| No. (%) (N=7,515)         | No. (%) (N=961)     | No. (%) (N=7,070)         |
| DOT                      |               |                           |                     |                           |
| HCW DOT                  | 115 (22)      | 1788 (24)                 | 184 (19)            | 1716 (24)                 |
| Family DOT               | 318 (62)      | 4407 (59)                 | 532 (55)            | 4193 (59)                 |
| SAT                      | 83 (16)       | 1323 (17)                 | 245 (26)            | 1161 (17)                 |
| Age                      |               |                           |                     |                           |
| 0–14                     | 11 (2)        | 118 (2)                   | 22 (2)              | 107 (1)                   |
| 15–44                    | 292 (57)      | 3913 (52)                 | 540 (56)            | 3665 (52)                 |
| 45–64                    | 114 (22)      | 2141 (28)                 | 235 (24)            | 2020 (29)                 |
| >65                      | 98 (19)       | 1336 (18)                 | 162 (17)            | 1272 (18)                 |
| Missing                  | 1 (0)         | 7 (0)                     | 2 (0)               | 6 (0)                     |
| Gender                   |               |                           |                     |                           |
| Male                     | 335 (65)      | 4966 (66)                 | 625 (65)            | 4676 (66)                 |
| Female                   | 180 (35)      | 2548 (34)                 | 335 (35)            | 2393 (34)                 |
| Missing                  | 1 (0)         | 1 (0)                     | 1 (0)               | 1 (0)                     |
| Nationality              |               |                           |                     |                           |
| Thai                     | 463 (90)      | 6895 (92)                 | 876 (91)            | 6482 (92)                 |
| Non-Thai                 | 52 (10)       | 611 (8)                   | 84 (9)              | 579 (8)                   |
| Missing                  | 1 (0)         | 9 (0)                     | 1 (0)               | 9 (0)                     |
| Marital status           |               |                           |                     |                           |
| Married                  | 282 (55)      | 4420 (59)                 | 506 (53)            | 4196 (59)                 |
| Non-Married              | 218 (42)      | 2822 (37)                 | 395 (41)            | 2645 (37)                 |
| Missing                  | 16 (3)        | 273 (4)                   | 60 (6)              | 229 (3)                   |
| Living in an urban district|           |                           |                     |                           |
| Urban district           | 226 (44)      | 3082 (41)                 | 462 (48)            | 2846 (40)                 |
| Non-urban district       | 275 (53)      | 4209 (56)                 | 459 (48)            | 4025 (57)                 |
| Missing                  | 15 (3)        | 224 (3)                   | 40 (4)              | 199 (3)                   |
| Mobile population*       |               |                           |                     |                           |
| Mobile                   | 211 (41)      | 1738 (23)                 | 385 (40)            | 1564 (22)                 |
| Non-mobile               | 284 (55)      | 5481 (73)                 | 514 (54)            | 5251 (74)                 |
| Missing                  | 21 (4)        | 296 (4)                   | 62 (6)              | 255 (4)                   |
| Ever treated with isoniazid preventive therapy (IPT) |           |                           |                     |                           |
| Previously treated with IPT | 2 (0)        | 55 (1)                    | 3 (0)               | 54 (1)                    |
| Not ever treated with IPT | 512 (100)    | 7406 (98)                 | 953 (99)            | 6965 (98)                 |
| Missing                  | 2 (0)         | 54 (1)                    | 5 (1)               | 51 (1)                    |
| Cough lasting >2 weeks at time of diagnosis |           |                           |                     |                           |
| Cough >2 weeks           | 314 (61)      | 5153 (68)                 | 576 (60)            | 4891 (69)                 |
| No cough >2 weeks        | 183 (35)      | 2010 (27)                 | 280 (29)            | 1913 (27)                 |
| Missing                  | 19 (4)        | 352 (5)                   | 105 (11)            | 266 (4)                   |
| Ever used injection drugs |           |                           |                     |                           |
| Ever used injection drugs | 21 (4)        | 183 (2)                   | 36 (4)              | 168 (2)                   |
| Not ever used injection drugs | 462 (90)  | 6631 (88)                 | 766 (80)            | 6372 (90)                 |
| Missing                  | 33 (6)        | 701 (9)                   | 159 (18)            | 575 (8)                   |
| In jail or prison         |               |                           |                     |                           |
| Previously in jail or prison | 12 (2)    | 116 (2)                   | 23 (2)              | 105 (1)                   |
| Not in jail/prison        | 489 (95)      | 6993 (93)                 | 825 (86)            | 6657 (94)                 |
| Missing                  | 15 (3)        | 406 (5)                   | 113 (12)            | 308 (5)                   |
| Diabetes mellitus         |               |                           |                     |                           |
**Table 2. cont.**

|                          | At Two Months |          |          | At End of Treatment |          |
|--------------------------|--------------|----------|----------|---------------------|----------|
|                          | Excluded     | Included | Excluded | Included            |          |
|                          | No. (%)      | No. (%)  | No. (%)  | No. (%)             |          |
|                          | (N = 516)    | (N = 7,515) | (N = 961) | (N = 7,070)         |          |
| Diabetes                 | 15 (3)       | 321 (4)  | 40 (4)   | 296 (4)             |          |
| No diabetes              | 414 (80)     | 5812 (77)| 710 (74) | 5516 (78)           |          |
| Missing                  | 87 (17)      | 1382 (18)| 211 (22) | 1258 (18)           |          |
| HIV status               |              |          |          |                     |          |
| Positive                 | 152 (30)     | 1564 (21)| 260 (27) | 1456 (20)           |          |
| Negative                 | 171 (33)     | 1623 (22)| 372 (39) | 4149 (59)           |          |
| Missing                  | 193 (37)     | 4328 (57)| 329 (34) | 1465 (21)           |          |
| Chest radiograph         |              |          |          |                     |          |
| Abnormal                 | 444 (86)     | 6746 (90)| 836 (87) | 6354 (90)           |          |
| Normal                   | 20 (4)       | 105 (1)  | 30 (3)   | 95 (1)              |          |
| Not performed or results missing | 52 (10) | 664 (9)  | 95 (10)  | 621 (9)             |          |
| Sputum culture result\(\#\) |            |          |          |                     |          |
| Growth of MTB            | 135 (26)     | 3629 (48)| 319 (33) | 3445 (49)           |          |
| No growth                | 88 (17)      | 1125 (15)| 141 (15) | 1072 (15)           |          |
| Not performed, contaminated, or grew NTM | 293 (57) | 2761 (37)| 501 (52) | 2553 (36)           |          |
| Initial treatment prescribed |        |          |          |                     |          |
| CAT I (2HRZE/4HR)        | 463 (90)     | 6902 (92)| 821 (85) | 6544 (93)           |          |
| Other regimens           | 53 (10)      | 613 (8)  | 140 (15) | 526 (7)             |          |
| Period of the year       |              |          |          |                     |          |
| Oct.–Dec.                | 98 (19)      | 1678 (22)| 145 (15) | 1631 (23)           |          |
| Jan.–Mar.                | 125 (24)     | 2011 (27)| 188 (20) | 1948 (28)           |          |
| Apr.–Jun.                | 138 (27)     | 2023 (27)| 235 (24) | 1926 (27)           |          |
| Jul.–Sep.                | 155 (30)     | 1802 (24)| 392 (41) | 1562 (22)           |          |
| Missing                  | 0 (0)        | 1 (0)    | 1 (0)    | 0 (0)               |          |

*Mobile was defined as not living in the same district for at least three of the past six months.
\(\#\) MTB denotes *Mycobacterium tuberculosis*, and NTM denotes non-tuberculous mycobacteria.
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**Discussion**

In this large, prospective observational study, we found that at least two months of DOT was associated with improved TB treatment outcomes. Although observation by either a HCW or family member was beneficial, the greatest magnitude of benefit was associated with HCW DOT and the greatest impact was on treatment default rates.

This is the largest analytical study every published about the impact of DOT on TB treatment outcomes. A major strength is that it was conducted among a diverse patient population within the existing public health care system in a high burden TB country. Previous studies of DOT have been conducted at specialized centers or, when community based, involved substantially smaller or more homogenous populations.[6] In this study, over 20% of patients were HIV-infected, and patients from the private sector, urban and rural districts, and migrant (non-Thai) populations were studied. We applied rigorous statistical techniques to control for the propensity of patients to receive DOT, adjusting for the fact that patients more likely to be adherent may also be more likely to consent to DOT and that DOT may be a marker for a healthcare facility with a better performing TB treatment program.

We found that having a treatment observer was better than not having a treatment observer in reducing default. Our findings were internally consistent. Because we only recorded whether DOT was provided for the first two months of treatment, we found that the impact of DOT was greatest on default at two months. Similarly, we found a gradient of impact for DOT, with HCW DOT having a larger impact than family DOT, a finding we would have expected given that HCWs are more likely to apply DOT strictly.[17] We found less benefit when we analyzed composite endpoints of default plus death, failure, or both. We would not expect DOT to have a substantial impact on death, because the primary risk factor for death during TB treatment in Thailand is HIV, and anti-retroviral therapy is the strongest determinant of survival in HIV-associated TB.[18] We also would not expect DOT to have a substantial impact on treatment failure, because rates of failure are low in Thailand, and, when it occurs, failure is likely attributable to drug resistance.[19] Our findings are also externally consistent with previous, smaller studies conducted in Thailand and in other settings on the benefit, albeit small, of DOT.[20–23] DOT by family members has also been shown to produce similar outcomes as HCW DOT in one randomized trial.[24]
Table 3. Bivariate and multivariate measures of association for successful TB treatment and health care worker observed, family member observed, and self-administered therapy.

| Outcomes | Treatment observer type | Analysis method |
|----------|-------------------------|-----------------|
|          | No. patients analyzed   | Health care worker | Family member | Self-administered | Bivariate | Propensity Score Risk Adjustment |
| On treatment vs. death or default at two months | HCW vs. SAT | 3100 | 1601/1784 (90%) | — | 1096/1316 (83%) | 1.8 (1.4–2.2) | 1.3 (1.0–1.7) |
| | Family vs. SAT | 5715 | — | 3782/4399 (86%) | 1096/1316 (83%) | 1.2 (1.0–1.5) | 1.1 (0.9–1.4) |
| | HCW vs. Family | 6183 | 1601/1784 (90%) | 3782/4399 (86%) | — | 1.4 (1.2–1.7) | 1.1 (0.9–1.3) |
| On treatment vs. default at two months | HCW vs. SAT | 2904 | 1601/1636 (98%) | — | 1096/1268 (86%) | 7.2 (4.9–10.4) | 3.8 (2.4–6.0) |
| | Family vs. SAT | 5255 | — | 3782/3987 (95%) | 1096/1268 (86%) | 2.9 (2.3–3.6) | 2.0 (1.5–2.7) |
| | HCW vs. Family | 5623 | 1601/1636 (98%) | 3782/3987 (95%) | — | 2.5 (1.7–3.6) | 2.1 (1.4–3.1) |
| Success vs. death, default, or failure at end of TB treatment | HCW vs. SAT | 2870 | 1369/1716 (80%) | — | 744/1154 (64%) | 2.2 (1.8–2.6) | 1.6 (1.3–2.0) |
| | Family vs. SAT | 5340 | — | 3130/4186 (75%) | 744/1154 (64%) | 1.6 (1.3–1.5) | 1.3 (1.1–1.5) |
| | HCW vs. Family | 5902 | 1369/1716 (80%) | 3130/4186 (75%) | — | 1.3 (1.2–1.5) | 1.1 (0.9–1.2) |
| Success vs. default at end of TB treatment | HCW vs. SAT | 2551 | 1369/1477 (93%) | — | 744/1074 (69%) | 5.6 (4.5–7.1) | 3.3 (2.4–4.5) |
| | Family vs. SAT | 4603 | — | 3130/3529 (89%) | 744/1074 (69%) | 3.5 (3.0–4.1) | 2.0 (1.6–2.4) |
| | HCW vs. Family | 4998 | 1369/1477 (93%) | 3130/3529 (89%) | — | 1.6 (1.3–2.0) | 1.5 (1.2–1.9) |

1HCW denotes health care worker directly observed therapy; family denotes family member directly observed therapy, and SAT denotes self-administered treatment (i.e., no directly observed therapy).

2For outcomes at two months, patients “on treatment” are considered successfully treated.
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Our study is subject to important limitations. First, this study was conducted within the routine healthcare system. We did not independently verify that patients recorded as receiving DOT actually received DOT, nor did we measure the rigor with which DOT was applied. Misclassification would most likely have involved patients being recorded as receiving DOT but not actually receiving it, which would bias our findings to a null association.[17] Therefore, we think that this study presents a conservative estimate of the benefit to DOT. Second, because of the large number of analyses performed, we could not perfectly balance all covariates when constructing propensity score quintiles. Within a given quintile, all patients should be equally likely to receive DOT (i.e., covariates should not be statistically associated with receiving DOT within that quintile); and, across all quintiles, less than 5% of covariates should be imbalanced.[15] In our analysis, we occasionally found imbalances of greater than 5% but less than 10%. We, therefore, also conducted traditional risk factor adjusted logistic regression for all 12 analyses, and found similar direction, magnitude, statistical significance, and precision as the propensity score analysis (data not shown). Although both statistical methods are valid, they can only adjust for measured confounders; it is possible that unmeasured confounders are responsible for the association between DOT and favorable outcomes that we found. Third, we only collected data about DOT use for the first two months of TB treatment. We are not able to draw any conclusions about the impact, either positive or negative, that providing DOT for the entire duration of treatment might have. Finally, we excluded patients from the two month and end of treatment analyses because of missing data. In some TB programs, patients who transfer out but who do not have final treatment outcomes reported as part of the original cohort are considered to have defaulted. We believe that either counting these patients as defaulters or excluding them would not change our findings, because the proportion of eligible patients excluded because of being transferred out was not substantial (359 [4.5%] for the two month outcome; 466 [5.8%] for the end-of-treatment outcome).

Despite the additional public health infrastructure provided through this project, treatment outcomes remained far below international targets. Some of this can be explained by the large number of HIV-infected, sputum smear-negative, private practice, and non-Thai patients included in our analysis. Nevertheless, Thailand’s overall treatment success rates remain sub-optimal. Through this project, treatment outcomes remained far below international targets. Some of this can be explained by the large number of HIV-infected, sputum smear-negative, private practice, and non-Thai patients included in our analysis. Nevertheless, Thailand’s overall treatment success rates remain sub-optimal. Despite the additional public health infrastructure provided through this project, treatment outcomes remained far below international targets. Some of this can be explained by the large number of HIV-infected, sputum smear-negative, private practice, and non-Thai patients included in our analysis. Nevertheless, Thailand’s overall treatment success rates remain sub-optimal.
program should strengthen its use of DOT and, wherever possible, use HCWs to provide it. Because a large number of patients that received SAT were treated in the private sector, efforts are also needed to bring private sector practices in line with international standards, including use of DOT.\[5\]

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Author Contributions

Conceived and designed the experiments: AA PL SN WW SK SM JKV. Performed the experiments: AA PL SN WW SK SM NG SKa PS JKV. Analyzed the data: AA PL SN WW SK SM NG SKa PS US PM JKV. Contributed reagents/materials/analysis tools: AA PS US PM JKV. Wrote the paper: AA JKV.