Analyzing of Tuberculosis in Turkey through SIR, SEIR and BSEIR Mathematical Models

Yasin Ucakan a,*, Seda Gulen b,b* and Kevser Koklu c,b

a Institute of Science, Mathematical Engineering, Davutpasa Campus, Yildiz Technical University, Istanbul, Turkey; b Faculty of Science and Art, Department of Mathematics, Tekirdag Namik Kemal University, Tekirdag, Turkey; c Yildiz Technical University, Faculty of Chemistry and Metallurgy, Department of Mathematical Engineering, Davutpasa Campus, Istanbul, Turkey

ABSTRACT
Since mathematical models play a key role in investigating the dynamics of infectious diseases, many mathematical models for these diseases are developed. In this paper, it is aimed to determine the dynamics of Tuberculosis (TB) in Turkey, how much it will affect the future and the impact of vaccine therapy on the disease. For this purpose, three mathematical models (SIR, SEIR and BSEIR) in the literature are considered for the case of Turkey. The model parameters are obtained with TB reported data from 2005 to 2015 by using the least square method. The obtained results revealed that the basic reproduction ratio for all three models is less than 1. Moreover, the stability analysis of the models and sensitivity analysis of the model parameters are presented and discussed. Finally, the accuracy of results for all three models is compared and the effect of the vaccination rate is discussed.

1. Introduction
During human history, many epidemics such as tuberculosis, influenza, SARS, MERS and Ebola have affected the population in many respect such as health, politics and economy. Therefore, scientists and governments have tried to keep epidemics under control. Recently, the current outbreak of the coronavirus disease (COVID-19) re-exposes the importance of epidemic researches and development of the mathematical models to describe the behaviour of epidemics. Among epidemic diseases, Tuberculosis (TB) which is a chronic infectious disease caused by Mycobacterium (MTB) usually affects the lungs but can also other organs like the brain, kidneys, gastrointestinal tract, bone, lymph nodes, etc. The bacteria that cause TB are spread through contaminated air released during the coughing of TB patients. Compared with other diseases caused by a single infectious agent, TB is the second leading lethal disease all over the world, especially in Asia and Africa. According to the World Health Organization (WHO) data, 1.8 million people died of the disease and 10.4 million fell ill in 2015. This shows that TB is a danger to human health and affects economic and social life...
negatively. Therefore, the current state of the disease should be understood and control programmes should be planned in order to prevent the spread of the disease.

Mathematical models have an important role in the planning of TB control programs. Modelling is a useful tool to understand the dynamics of an epidemic that would help to prevent spreading. Besides, models contribute to predict the future of epidemic and control of the disease. The first study on the mathematical modelling of the spread of disease was proposed by Bernoulli in 1766 [1]. Since the 20th century, interest in studying mathematical modelling of infectious diseases has increased. The earlier related studies can be found in [2–9].

In 1927, Kermack and McKendrick proposed a deterministic model to describe the behaviour of epidemic propagation known as the Susceptible – Infected – Recovered (SIR) model [10]. Although this model has been successfully used to describe the behaviour of disease, neglecting the other compartments and control strategies, such as vaccination, treatment, quarantine, isolation, and the effect of age and sex is unrealistic. Therefore, many researchers have focused on constructing more realistic models [11–24].

A significant improvement to the SIR model is the addition of the exposed group which is infected but not infectious, called the Susceptible – Exposed – Infected – Recovered (SEIR) model [25–29]. The role of the season on the transmission of an epidemic was first investigated by Aron and Schwartz [30] using the SEIR model. Li et al. [31] studied the global dynamics of the SEIR model in the case of variable total population size. Newton and Reiter [32] developed an SEIR model for observing the behaviour of dengue fever. After that, the studies on TB began to model by the SEIR models. Chavez and Feng [33] focused on four models to understand the disease transmission dynamics of TB. Röst and Wu [34] proposed a new SEIR model in which the infectivity depends on age. Dontwi et al. [35] described the spreading of TB in Amansie West District Ghana by using the standard SEIR model. Yali Yang et al. [36] evaluated the cost of control strategies by using an SEIR model. Side et al. [37] proposed a SIR and an SEIR models for TB and analysed these models. Zhang et al. [38] set up a new mathematical model for TB in China using the data from January 2005 to December 2012. Xu et al. [39] proposed a mathematical model to investigate the control and precautions in Guangdong of China. Besides, many authors interested in global stability of TB models [40–43].

One of the most important factors in preventing and controlling the spread of tuberculosis is vaccination. Since 1921, the Bacillus Calmette-Guérin (BCG) vaccine remains the most widely used vaccine for the prevention of TB. Evidence has shown that BCG has a protective efficacy of about 75% in preventing some serious types of TB (e.g. meningitis) in children [44]. BCG is currently administered to newborns of high-risk populations as part of the World Health Organization (WHO) Expanded Programme on Immunization (EPI) [45]. Therefore, mathematical modelling of TB including vaccination has gained more importance to make more accurate predictions. In literature, there are many studies on this topic for various epidemics [46–50]. Specifically for TB, Liu et al. [51] proposed a mixed vaccination strategy that is the combination of constant vaccination and pulse vaccination. Egbetade and Ibrahim [52] set up a new mathematical model incorporating treatment, migration and vaccination. Rangkuti et al. [53] explained the spread of TB in North Sumatera Indonesia using VSEIR, which was created by adding the vaccination parameter to the SEIR model. Egonmwan et al. [54] formulated a mathematical model that
incorporates vaccination of newborn children and older susceptible individuals into the transmission dynamics of TB in a population.

Most of the epidemic diseases are modelled by a system of nonlinear ordinary differential equations with respect to model parameters and state variables. The main problem in these models is to determine the model parameters that describe the behaviour of the model in a realistic way. Generally, these parameters are adjusted by using nonlinear optimization techniques. Obtaining the parameters is important to calculate the basic reproduction number, $R_0$, which represents the expected number of new cases of infectious generated by an infected individual. $R_0$ is a key to understand how fast the disease will spread and the impact of control strategies. If $R_0 > 1$, disease breaks out into epidemics, but $R_0 < 1$, disease dies out.

TB is a lethal disease and is still struggled hard to control for some countries. Even though TB is controlled in Turkey, the geopolitical position of the country and letting in too many immigrants always has put it at risk. Therefore, the development of researches and prevention strategies should be continued. Although the studies [55,56] have been successfully used to describe the spreading of TB in Turkey, in these models, the assumption of absence of some factors such as the birth and the death rate, exposed individuals and prevention, lead to unrealistic estimations. Therefore, in this paper, TB disease in Turkey has been analysed by three epidemiological models (a modified SIR, an SEIR, and a BSEIR) to obtain more realistic predictions. Our aim is to investigate and discuss the characteristics of all three models regarding TB in Turkey, the information they provide and the situations in which the models are used. Besides, one of our main interests is remarking the effect of vaccination on the propagation of virus. In the modified SIR model, the population has been characterized by a death rate and a birth rate equal to the death rate. Through using the SEIR model, it has been included a latent or incubation time which means a certain time for an infected individual to become infectious. The control of the spread of TB through BCG vaccination therapy has been investigated with the BSEIR model. Besides, the model parameters have been determined by fitting the real data reported by the WHO [57–67]. The obtained results concluded that the basic reproduction number for all three models is $R_0 < 1$. This means that the disease in Turkey is not alarming, but since the number of patients does not decrease dramatically, the control strategies should be continued. The stability analysis of the disease-free equilibrium and endemic equilibrium points is investigated for all three models. Moreover, the sensitivity analysis of the model parameters has been performed and their simulation results have been plotted. In addition, the 95% confidence interval estimate graphs have been presented. The computations show that the predictions produced by all three models approximate the real data very well. Besides, it is emphasized that the model including the vaccination factor (the BSEIR) presents more realistic approaches. Furthermore, based on our literature review, such a comprehensive study is not worked through TB for Turkey so far.

The paper is organized as follows: In Section 2, the models, the modified SIR, the SEIR, and the BSEIR, for describing the dynamics of Tuberculosis are introduced and the equilibria and stability analysis of the models are studied. In addition, the sensitivity analysis of the parameters of the models is investigated. Then, the numerical simulations of all three models are presented and discussed how this is fitted against the data available in Turkey in Section 3. Besides, the predictions of the three models are compared and the effect of vaccination is remarked. Finally, it is concluded the paper in Section 4.
2 Material and methods

2.1. Mathematical modelling

2.1.1. The modified SIR model

Since the basic SIR model proposed by Kermack and McKendrick neglects any natural deaths and births, some authors modified the model by considering a population by a death and a birth rates. Thus, the nonlinear system of differential equations including the death and the birth rates for TB is given by

\[
\begin{align*}
\frac{dS(t)}{dt} &= bN - \beta S(t)I(t)/N - \mu S(t) \\
\frac{dI(t)}{dt} &= \beta S(t)I(t)/N - (\gamma + \mu)I(t) \\
\frac{dR(t)}{dt} &= \gamma I(t) - \mu R(t)
\end{align*}
\]

with subject to

\[S(0) \geq 0, \quad I(0) \geq 0 \quad \text{and} \quad R(0) \geq 0,\]

where \(\beta, \gamma, \mu,\) and \(b\) are the transmission rate, the recovery rate, the birth rate and the death rate, respectively. In this model, it is assumed that the death and the birth rates are equal to each other. This model comprises three subgroups: \(S(t)\) represents the number of non-infected but liable to infection in the population at time \(t, I(t)\) is the number of the infected individuals who can transmit the disease to the susceptible individuals in the population and \(R(t)\) is the recovered individuals who have been immunized and they are not able to infect again. The total population which is homogeneous and isolated is denoted by \(N = S(t) + I(t) + R(t)\) for all \(t\). The transmission coefficient \(\beta\) estimates the probability of getting the disease in contact between a susceptible and an infectious individuals. This model does not take into account age, vaccination or waning immunity. Since \(\beta\) and \(\gamma\) are the transition rate, are stated as probabilities, their range is \(0 < \beta, \gamma < 1\).

Equilibrium points and stability. The dimensionless form of the system (1) is presented by

\[
\begin{align*}
\frac{dS}{dt} &= -\beta S(t)I(t) + \mu(1 - S(t)) \\
\frac{dI}{dt} &= \beta S(t)I(t) - (\gamma + \mu)I(t)
\end{align*}
\]

where \(R(t) = 1 - S(t) - I(t)\). The third equation in system (1) can be omitted because the first two equations do not depend on it. To find the equilibrium points, the system (2) should be equated to zero:

\[
\frac{dS(t)}{dt} = \frac{dI(t)}{dt} = 0
\]

Then the equilibrium points are obtained, namely disease-free equilibrium (DFE) and endemic equilibrium (EE), respectively:
\[ D_1 = (S_1^*, I_1^*) = (1, 0), \quad \text{and} \quad D_2 = (S_2^*, I_2^*) = \left( \frac{\gamma + \mu}{\beta}, \frac{\mu(\beta - \gamma - \mu)}{\beta(\mu + \gamma)} \right). \] (3)

The Jacobian matrices for the system (2) are evaluated at the DFE and the EE as follows:

\[
J(D_1) = J(S_1^*, I_1^*) = \begin{bmatrix}
-\mu & \beta \\
0 & \beta - \gamma - \mu
\end{bmatrix},
\]

\[
J(D_2) = J(S_2^*, I_2^*) = \begin{bmatrix}
-\frac{\mu\beta}{\mu + \gamma} & \frac{\mu(\beta - \gamma - \mu)}{\mu + \gamma} \\
-\gamma - \mu & 0
\end{bmatrix}.
\]

In order to the equilibrium points of the system (2) are stable, the eigenvalues must be negative or have a negative real part [68]. The eigenvalues can be obtained by solving the characteristic equation \( \lambda^2 - Tr(J)\lambda + det(J) = 0 \). Since the eigenvalues must satisfy the rule stated above, \( det(J) > 0 \) and \( Tr(J) < 0 \) must be proven.

\[ det(J_{D_1}) = -\mu(\beta - \gamma - \mu) > 0, \]

and

\[ Tr(J_{D_1}) = \beta - \gamma - 2\mu < 0, \quad \mu > 0. \]

Hence, \( \frac{\beta}{\gamma + \mu} < 1 \).

\[ det(J_{D_2}) = \mu(\beta - \gamma - \mu) > 0, \]

and

\[ Tr(J_{D_2}) = \frac{-\mu\beta}{\mu + \gamma} < 0, \quad \mu > 0. \]

Hence, \( \frac{\beta}{\gamma + \mu} > 1 \).

This includes:

The DFE \( D_1 \) is locally asymptotically stable if and only if \( \frac{\beta}{\gamma + \mu} < 1 \), otherwise unstable.

The EE \( D_2 \) is locally asymptotically stable if and only if \( \frac{\beta}{\gamma + \mu} > 1 \), otherwise unstable.

Here, \( \frac{\beta}{\gamma + \mu} \) is a threshold value, called the basic reproduction number \( R_0 \), which has an important role in determining dynamics of the disease and is used for prevention strategies. It helps to decide whether the disease spreads or eradicates. If the basic reproduction number is less than 1, then the disease will not spread in the population and dies out. Otherwise, the disease breaks out into an epidemic. Therefore, \( R_0 \) is a crucial significance in mathematical and biological aspects.

### 2.1.2. The SEIR model

The SEIR epidemic model is a generalization of the SIR model. This model on the spread of TB consists of four compartments, namely susceptible \((S(t))\), Exposed \((E(t))\), Infected \((I(t))\) and Recovered \((R(t))\), can be interpreted as follows:
\[
\begin{align*}
\frac{dS(t)}{dt} &= bN - \beta S(t)I(t)/N - \mu S(t) \\
\frac{dE(t)}{dt} &= \beta S(t)I(t)/N - (\varepsilon + \mu)E(t) \\
\frac{dI(t)}{dt} &= \varepsilon E(t) - (\gamma + \mu)I(t) \\
\frac{dR(t)}{dt} &= \gamma I(t) - \mu R(t)
\end{align*}
\]

(4)

with subject to

\[
S(0) \geq 0, \quad E(0) \geq 0, \quad I(0) \geq 0, \quad \text{and} \quad R(0) \geq 0,
\]

where a new compartment \(E(t)\) which denotes the individuals who are infected but the symptoms of the disease are not yet visible and the parameter \(\varepsilon\) is the rate at which the exposed individuals become infective so that \(\frac{1}{\varepsilon}\) is the mean latent period. The transmission rate, the recovery rate, the birth and death rates are represented by \(\beta, \gamma, \mu\) and \(b\), respectively. This model assumes that the death and the birth rates are equal to each other. The total population which is homogeneous and isolated is denoted by \(N = S(t) + E(t) + I(t) + R(t)\) for all \(t\). In this model, the other factors such as age, sex, vaccination, etc. are neglected.

**Equilibrium points and stability.** The dimensionless form of the system (4) is presented by

\[
\begin{align*}
S' &= -\beta S(t)I(t) + \mu(1 - S(t)) \\
E' &= \beta S(t)I(t) - (\varepsilon + \mu)E(t) \\
I' &= \varepsilon E(t) - (\gamma + \mu)I(t)
\end{align*}
\]

(5)

where \(R(t) = 1 - S(t) - I(t) - E(t)\). The fourth equation in system (4) can be omitted because the first three equations do not depend on it. To find the equilibrium points, the system (5) should be equated to zero:

\[
\frac{dS(t)}{dt} = \frac{dE(t)}{dt} = \frac{dI(t)}{dt} = 0
\]

Then the equilibrium points are obtained, namely DFE and EE, respectively:

\[
\begin{align*}
\text{D1} &= (S_1^*, E_1^*, I_1^*) = (1, 0, 0), \\
\text{D2} &= (S_2^*, E_2^*, I_2^*) = \left( \frac{1}{R_0}, \frac{\mu(R_0 - 1)}{R_0(\varepsilon + \mu)}, \frac{\mu(R_0 - 1)}{\beta} \right).
\end{align*}
\]

(6)

Here, \(R_0 = \frac{\beta \varepsilon}{(\gamma + \mu)(\varepsilon + \mu)}\) is the basic reproduction number [69]. The threshold quantity \((R_0)\) can be interpreted as the product of the contact rate \((\beta)\) and average fraction \(\left(\frac{\varepsilon}{\varepsilon + \mu}\right)\) surviving the incubation period and average infectious period \(\left(\frac{1}{\gamma + \mu}\right)\) [35]. The Jacobian matrices for the system (5) are evaluated at the DFE and the EE as follows:
\[ J(D_1) = J(S_1^*, E_1^*, I_1^*) = \begin{bmatrix} -\mu & 0 & -\beta \\ 0 & -\mu - \epsilon & -\beta \\ 0 & \epsilon & -\mu - \gamma \end{bmatrix}, \]

\[ J(D_2) = J(S_2^*, E_2^*, I_2^*) = \begin{bmatrix} -\mu R_0 & 0 & -\beta R_0 \\ \mu (R_0 - 1) & -\mu - \epsilon & -\beta R_0 \\ 0 & \epsilon & -\mu - \gamma \end{bmatrix}. \]

In order to the equilibrium points of the system (5) are stable, the eigenvalues must be negative or have a negative real part \([68]\). The eigenvalues can be obtained by solving the characteristic equation corresponding to \(J(D_1)\) and \(J(D_2)\).

**i. Stability analysis of DFE** Characteristic equation for \(D_1\) is

\[ (\mu + \lambda)[\lambda^2 + (y + 2\mu + \epsilon)\lambda + (\mu + \epsilon)(\mu + \gamma)(1 - R_0)] = 0. \tag{7} \]

Then the eigenvalues are

\[ \lambda_1 = -\mu, \]

\[ \lambda_2 = \frac{1}{2}[-(y + 2\mu + \epsilon) - \sqrt{(y + 2\mu + \epsilon)^2 - 4(\mu + \epsilon)(\mu + \gamma)(1 - R_0)}], \tag{8} \]

\[ \lambda_3 = \frac{1}{2}[-(y + 2\mu + \epsilon) + \sqrt{(y + 2\mu + \epsilon)^2 - 4(\mu + \epsilon)(\mu + \gamma)(1 - R_0)}]. \]

\(\lambda_1\) and \(\lambda_2\) are negative, clearly. \(\lambda_3 < 0\) only for \(R_0\) less than 1. Hence, \(D_1\) is locally asymptotically stable if and only if \(R_0 < 1\), otherwise unstable.

**ii. Stability analysis of EE**

Characteristic equation for \(D_2\) is

\[ \lambda^3 + (\epsilon + y + \mu(2 + R_0))\lambda^2 + \mu R_0(\epsilon + y + 2\mu)\lambda + \mu(\mu + \epsilon)(\mu + \gamma)(R_0 - 1) = 0. \tag{9} \]

Methods that allow determining whether all roots have negative real parts and ensuring the stability of the system without solving the characteristic equation are of great importance. The Routh-Hurwitz criterion, which contains the necessary and sufficient conditions for the stability of the system, is one of these methods.

**Theorem 2.1. (Routh-Hurwitz Stability Criterion)** \([70]\)

*Given the polynomial,*

\[ P(\lambda) = \lambda^n + a_1\lambda^{n-1} + \ldots + a_{n-1}\lambda + a_n, \tag{10} \]

*where the coefficients \(a_i\) are the real constants, \(i = 1, \ldots, n\), define the \(n\) Hurwitz matrices using the coefficient \(a_i\) of the characteristics polynomial:*
H_1 = [a_1], H_2 = \begin{bmatrix} a_1 & 1 \\ a_3 & a_2 \end{bmatrix}, H_3 = \begin{bmatrix} a_1 & 1 & 0 \\ a_3 & a_2 & a_1 \\ a_5 & a_4 & a_3 \end{bmatrix} \text{ and } H_n = \begin{bmatrix} a_1 & 1 & 0 & \cdots & 0 \\ a_3 & a_2 & a_1 & \cdots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \cdots & a_n \end{bmatrix}

where a_j = 0 if j > n. All the roots of the polynomial P(\lambda) are negative or have negative real part if and only if the determinants of all Hurwitz matrices are positive;

\[ \det H_j > 0, \quad j = 1, 2, \ldots, n. \]

The Routh-Hurwitz criteria for which n = 3 taken can be simplified as follows

\[ \det H_1 = a_1 > 0 \]

\[ \det H_2 = \det \begin{bmatrix} a_1 & 1 \\ a_3 & a_2 \end{bmatrix} = a_1 a_2 - a_3 > 0 \]

\[ \det H_3 = \det \begin{bmatrix} a_1 & 1 & 0 \\ a_3 & a_2 & a_1 \\ 0 & 0 & a_3 \end{bmatrix} = a_3 (a_1 a_2 - a_3) > 0, \quad (a_3 > 0). \]

From characteristic equation (9), \( a_1 = \varepsilon + \gamma + \mu(2 + R_0) > 0 \) is provided. It is clear that \( a_1 a_2 - a_3 > 0 \) for values \( a_2 = \mu R_0 (\varepsilon + \gamma + 2 \mu) \) and \( a_3 = \mu (\mu + \varepsilon)(\mu + \gamma)(R_0 - 1) \). Hence, \( a_3 > 0 \) if and only if \( R_0 > 1 \). Hence, it can be concluded that:

The EE D_2 is locally asymptotically stable if and only if \( R_0 > 1 \), otherwise unstable.

2.1.3. The BSEIR model

In this subsection, to investigate the effect of vaccination on the behaviour of TB, BSEIR model proposed by Liu et al. [51] can be described by the dynamic system of differential equations as follows:

\[
\begin{aligned}
\frac{dB(t)}{dt} &= \Lambda p - kB(t) \\
\frac{dS(t)}{dt} &= kB(t) + \Lambda (1 - p) - \beta S(t) I(t)/N - \mu S(t) \\
\frac{dE(t)}{dt} &= \beta S(t) I(t)/N - (\varepsilon + \mu) E(t) \\
\frac{dI(t)}{dt} &= \varepsilon E(t) - (\gamma + \mu + d) I(t) \\
\frac{dR(t)}{dt} &= \gamma I(t) - \mu R(t)
\end{aligned}
\]  

(11)

with subject to

\[ B(0) \geq 0, \quad S(0) \geq 0, \quad E(0) \geq 0, \quad I(0) \geq 0 \quad \text{and} \quad R(0) \geq 0, \]

where the parameters \( p \) (0 < \( p < 1 \)) and \( \Lambda \) denote the fraction of the newborns vaccinated successfully and the recruitment rate, respectively. Since the positive effect of BCG vaccination is limited, the vaccinated successfully individuals become susceptible again by the rate \( k \). The transmission rate, the recovery rate, the natural death rate and the disease-induced death rate are represented by \( \beta \), \( \gamma \), \( \mu \) and \( d \), respectively. The model comprises five subgroups: BCG vaccinated \( B(t) \), Susceptible
$S(t)$, Exposed $E(t)$, Infected $I(t)$ and Recovered $R(t)$. In addition to the SEIR model, the BSEIR model includes the BCG vaccinated subgroup which denotes the vaccinated newborns successfully. In the BCG protection period, they will not get infected even if they contact infected individuals when the vaccination provides immunity to all of them. The total population, $N = B(t) + S(t) + E(t) + I(t) + R(t)$, changes with respect to time since the natural death rate and the birth rate do not have taken equally. Besides, since the vaccination is thought to prevent for only 10 to 15 years and the natural death of children is about %1, the natural death rate in subgroup $B(t)$ is neglected.

**Equilibrium points and stability.** To find the equilibrium points, the system (11) should be equated to zero:

$$
\frac{dB(t)}{dt} = \frac{dS(t)}{dt} = \frac{dE(t)}{dt} = \frac{dI(t)}{dt} = \frac{dR(t)}{dt} = 0
$$

Then the equilibrium points are obtained, namely DFE and EE, respectively:

$$
D_1 = \left( b_1^*, s_1^*, e_1^*, i_1^*, r_1^* \right) = \left( \frac{\Delta p}{k}, \Delta \mu, 0, 0, 0 \right),
$$

$$
D_2 = \left( b_2^*, s_2^*, e_2^*, i_2^*, r_2^* \right) = \left( \frac{\Delta p}{k}, \frac{\beta \Lambda s^2}{\beta \Lambda s^2 + N \mu}, \frac{\beta \Lambda s^2}{\beta \Lambda s^2 + N \mu}, \frac{\beta \Lambda s^2}{\beta \Lambda s^2 + N \mu}, i_2^*, r_2^* \right),
$$

where $i_2^* = \frac{\beta \in A-N \mu(\epsilon+\mu)(\mu+d+y)}{\beta(\epsilon+\mu)(\mu+d+y)}$. We can obtain the basic reproduction number $R_0$ by using the next-generation matrix [71]:

$$
R_0 = \rho(FV^{-1}) = \frac{\beta \in k}{(y+\mu+d)(\epsilon+\mu)(\mu p+k)},
$$

where $\rho$ denotes the spectral radius and the matrices $F$ and $V$ are given by

$$
F = \begin{bmatrix}
0 & \frac{\beta k}{\mu p+k} \\
0 & 0
\end{bmatrix},
V = \begin{bmatrix}
\epsilon + \mu & 0 \\
-\epsilon & \mu + d + y
\end{bmatrix}.
$$

The Jacobian matrix for the system (11) is evaluated at the EE as follows:

$$
J(D_2) = \begin{bmatrix}
-k & 0 & 0 & 0 & 0 \\
k & -\mu & 0 & -\frac{\beta \Lambda}{N \mu} & 0 \\
0 & 0 & -(\epsilon + \mu) & \frac{\beta \Lambda}{N \mu} & 0 \\
0 & 0 & \epsilon & -(y + \mu + d) & 0 \\
0 & 0 & 0 & \gamma & -\mu
\end{bmatrix}.
$$

Solving $det(J - \lambda I) = 0$, the characteristic equation is

$$
(-k - \lambda)(-\mu - \lambda)^2(\lambda^2 + a_1 \lambda + a_2) = 0
$$

where

$$
a_1 = \gamma + 2\mu + d + \epsilon, \\
a_2 = (\epsilon + \mu)(\gamma + \mu + d) - \frac{\epsilon \beta \Lambda}{N \mu}.
$$
Roots of the characteristic equations are $\lambda_1 = -k, \lambda_2 = -\mu$ and $\lambda_3 = -\mu$ are negative and other two roots satisfies the following quadratic equation

$$\lambda^2 + a_1\lambda + a_2 = 0.$$  \hspace{1cm} (14)

Clearly, if $a_1 > 0$ and $a_2 > 0$, two roots of (14) will be negative. Hence, $D_1$ is locally asymptotically stable if and only if $R_0 < 1$, otherwise unstable.

### 2.2. Numerical solution and estimation of model parameters

The models in the present work have been numerically solved using software MATLAB R2019b by the command 'ode15s'. The reason for choosing this command is the stiffness of differential equation systems that generate the SIR, the SEIR and the BSEIR models. Since stiff problems are characterized by significantly different magnitudes of variation rates, the command 'ode45' may struggle to solve the system.

The model parameters have been adjusted based on TB incidence data taken from the WHO Global Tuberculosis Report [57–67] from 2005 to 2015. Some of the parameters have been obtained from the literature [72,73], as seen in Table 1 and others have been assumed or fitted from data. The death rate $\mu$ has been estimated at approximately $\mu = \frac{366,471}{73,722,988} = 0.005$, where 366,471 is the number of death individuals and 73,722,988 is the population of Turkey in 2010 [73] as reference. The birth rate $b$ is 0.01737 [73] so the recruitment rate is $\Lambda = 1,280,806$ per year ($\Lambda = b \times N$). Other parameters have been adjusted through the minimization of the quadratic objective function

$$\phi = \sum_{i=1}^{n} (I_i^e - I_i^a)^2$$  \hspace{1cm} (15)

using least square method. Here, $I_i^a$, $I_i^e$ and $n$ denote the actual TB infected case, the model solution at time $t_i$ and the number of available data, respectively. To minimize the

| Parameters/Initial data | Description | Value a | Value b | Value c | Source |
|-------------------------|-------------|---------|---------|---------|--------|
| $\beta$                 | Rate of waning immunity | –       | –       | 0.054   | Fitted |
| $\gamma$                | Rate of progression to infectious stage from the exposed | –       | 1.428   | 1.435   | Fitted |
| $\mu$                   | Recovery rate | 0.939   | 0.935   | 0.935   | Fitted |
| $\phi$                  | Natural death rate | 0.0049  | 0.0049  | 3.0049  | [73]   |
| $\psi$                  | Disease-induced death rate | –       | 0.03    | 0.03    | [57]   |
| $S_0$                   | Initial number of susceptible | 67,610,005 | 67,595,153 | 63,095,153 | Assumed |
| $E_0$                   | Initial number of exposed | 14,852  | 14,852  | 14,852  | Assumed |
| $I_0$                   | Initial number of infected | 20,535  | 20,535  | 20,535  | [57]   |
| $R_0$                   | Initial number of recovered | 1,230,000| 1,230,000| 1,230,000| Assumed |
| $R_0$                   | Reproduction number | 0.9664  | 0.9211  | 0.9045  | Calculated |

\(^a\)SIR Model
\(^b\)SEIR Model
\(^c\)BSEIR Model
function (15) it has been used the command ‘nlinit’ which solves nonlinear regression problems through the Levenberg-Marquardt algorithm in MATLAB R2019b.

### 2.3. Sensitivity analysis

Sensitivity analysis provides to analyse the effect of each parameter on disease transmission and prevalence. It is commonly used to determine the robustness of model predictions to parameter values because of errors in data collection and assumed parameters. It is important to determine parameters that have a high impact on $R_0$ and should be targeted by intervention strategies. Therefore, here, the partial derivatives of the basic reproduction number $R_0$ with respect to model parameters $p$, $k$, $\beta$ and $\gamma$ have been calculated. Since $\frac{\partial R_0}{\partial k} > 0$, increasing $k$ means increasing $R_0$. It shows that the number of infected will increase faster. Parameter $p$ is the fraction of vaccinated successfully and $\frac{\partial R_0}{\partial p} < 0$, and parameter $\gamma$ is the recovery rate and $\frac{\partial R_0}{\partial \gamma} < 0$, also. Hence, it can be said that the total number of infected populations can be reduced by increasing the parameters $p$ and $\gamma$. Besides, being $\frac{\partial R_0}{\partial \beta} > 0$, it is meant that the infection can be reduced by decreasing the transmission rate $\beta$.

To estimate the relative change in a variable when parameters change, sensitivity indices should be calculated. The calculation of these indicates has been executed by means of the following definition.

**Definition.** The normalized forward sensitivity index of $R_0$, which is differentiable with respect to a given parameter $\sigma$, is defined by

$$S_0^\sigma = \frac{\sigma \frac{\partial R_0}{\partial \sigma}}{R_0} \tag{16}$$

[71].

The obtained sensitivity indices of the basic reproduction number $R_0$ for the baseline model parameters calculated by the formula (16) are presented in Table 3.

### 3. Results and discussion

In this section, to illustrate the numerical results for all three models, initial data have been taken as the following values. Since the epidemic data from 2005 to 2015 have been taken into account, the total initial population has been accepted as $N(0) = 68,860,540$ as the same as the reported population of Turkey in 2005 [72]. The initial infected population was given in the report [57] as $I(0) = 20,535$. $E(0)$ has been assumed by comparing the rate of the infected to exposed individuals in literature and adapting to data in Turkey. $R(0)$ has been assumed by taking into account recovered individuals by the rate of success of therapy before 2005, average life span and natural death rate. $B(0)$ has been estimated considering the number of newborns in recent years reported by the
For the initial susceptible populations for the SIR, the SEIR and the BSEIR are given by, respectively,

\[ S(0) = N - I(0) - R(0), \]

\[ I(0) = \frac{S(0)}{R(0)}, \]

\[ R(0) = 1. \]

**Figure 1.** The bar diagrams for the model predictions and reported infected data.

**Figure 2.** The behaviours of compartments population over time for the SIR model.
\[ S(0) = N - E(0) - I(0) - R(0), \]

\[ S(0) = N - B(0) - E(0) - I(0) - R(0). \]

As indicated in Section 2.2, the parameters \( \beta, \gamma, \epsilon, p, k \) are fitted from real data by using the minimization function (15). The fitted parameter values and initial data have been listed in Table 1.

The epidemic models generated by the systems (1), (4) and (11) have been solved by MATLAB R2019b using command ‘ode15s’. This command uses numerical backward differentiation formulas with a maximum order of \( k = 5 \) using a quasi constant step method. Since the timelines of real incidence data are according to years, the timelines of obtained numerical solutions have been taken by specifying the interval of integration as a vector of years in the ode15s.

(Figure 1) presents the bar diagram of the per year total infection of real cases and model predictions for all three models. The obtained results revealed that the model predictions approximate the real data very well.

In (Figures 2–4), it is plotted graphs to show the behaviours of all groups of individuals for all three models. The variation of \( S(t) \), \( I(t) \) and \( R(t) \) for the modified SIR model is displayed in (Figure 2). As seen in the figure, the population of infected with TB has decreased and hence, the populations of recovered have increased. With this increase, a decrease in the number of susceptible populations is observed. Therefore, it is concluded that the TB disease is under control. Also, the behaviour of the compartments

**Figure 3.** The behaviours of compartments population over time for the SEIR model.
of the SEIR model given by (Figure 3) is similar to the SIR model. Additionally, the exposed population has been decreasing since the infected individuals have been decreasing. However, if the absence of control strategies such as treatment, vaccination, etc., the infected individuals in the population will increase, and therefore, TB becomes an epidemic.

The behaviour of the compartments of the BSEIR model has been simulated in (Figure 4). Unlike the SIR and the SEIR models, the BSEIR model has included the vaccination rate, and the birth and the death rates are not equal to each other in this model. For the prevention of TB, the BCG vaccine is widely used, which is 80% effective in preventing TB and the duration is about 10 years [75]. Therefore, vaccination is one of the treatments for TB patients and hence, an assumption for the transfer of a proportion of the susceptible population to the vaccination class is considered. As can be seen in (Figure 4), the B(t) class is increasing. However, since the population is not constant, S(t) class shows an increase. The fact that population growth does not have a negative effect on the number of patients due to the vaccination factor. Since the high recovery rate and the effect of the vaccine, the number of infected decreases.

The behaviour of infected individuals for all three models is plotted in (Figure 5). As displayed in the figure, the reported real incidence data and the obtained predictions for all three models are in good agreement. From the figure, it can be seen the superiority of the BSEIR model when compared to the SIR and the SEIR models. The factors that make the BSEIR model more realistic are the active TB vaccine is included in the model and the population is not constant. Since BCG vaccination is used in Turkey for a long time, when taking into account the vaccination rate has been revealed more realistic...
predictions. Although the estimates obtained from the SIR and the SEIR models are close to the real data, it may not provide reliable solutions because of ignoring the vaccination.

To show accuracy of the numerical solutions of the SIR, the SEIR and the BSEIR, the relative errors are calculated by

\[ \text{Relative Error} = \frac{|\text{Model} - \text{Data}|}{\text{Data}} \]
Figure 7. The total number of infected individuals for different values of $\beta$ and $\gamma$ (SEIR model).

Table 2. Prediction of infected population for the SIR, the SEIR and the BSEIR.

| Year | Reported data | Numerical value a | Error | Numerical value b | Error | Numerical value c | Error |
|------|---------------|-------------------|-------|-------------------|-------|-------------------|-------|
| 2005 | 20535         | 20535             | –     | 20535             | –     | 20535             | –     |
| 2006 | 20526         | 19621             | 0.04410 | 20543             | 0.00085 | 20693             | 0.00816 |
| 2007 | 19694         | 18743             | 0.04830 | 19537             | 0.00797 | 19594             | 0.00507 |
| 2008 | 18452         | 17896             | 0.03014 | 18486             | 0.00183 | 18456             | 0.00021 |
| 2009 | 17402         | 17084             | 0.01825 | 17477             | 0.00432 | 17389             | 0.00075 |
| 2010 | 16551         | 16306             | 0.01479 | 16522             | 0.00177 | 16404             | 0.00886 |
| 2011 | 15679         | 15561             | 0.00750 | 15618             | 0.00390 | 15501             | 0.01134 |
| 2012 | 14691         | 14848             | 0.01071 | 14762             | 0.00483 | 14677             | 0.00095 |
| 2013 | 13409         | 14167             | 0.05655 | 13952             | 0.04052 | 13928             | 0.03881 |
| 2014 | 13378         | 13516             | 0.01031 | 13187             | 0.01427 | 13253             | 0.00917 |
| 2015 | 12772         | 12894             | 0.00956 | 12463             | 0.02421 | 12649             | 0.00949 |
| 2016 | 12417         | 12300             | 0.00942 | 11779             | 0.05138 | 12111             | 0.02464 |
| 2017 | 12046         | 11733             | 0.02598 | 11132             | 0.07588 | 11635             | 0.03412 |
| 2018 | 11786         | 11192             | 0.05040 | 10520             | 0.10742 | 11220             | 0.04802 |
| 2019 | –             | 10675             | –     | 9942              | –     | 10861             | –     |
| 2020 | –             | 10182             | –     | 9396              | –     | 10558             | –     |
| 2021 | –             | 9713              | –     | 8679              | –     | 10306             | –     |
| 2022 | –             | 9264              | –     | 8391              | –     | 10105             | –     |
| 2023 | –             | 8836              | –     | 7930              | –     | 9952              | –     |

aSIR Model  
bSEIR Model  
cBSEIR Model

\[ h(t) = \frac{|I_e(t) - I_d(t)|}{I_d(t)}, \]  

(17)

where $I_e(t)$ and the $I_d(t)$ are the model prediction and the corresponding data at time $t$, respectively. The reported infected data, model predictions and relative errors have been listed in (Table 3). In the calculation, it is used the reported infected data from 2005 to 2015 to compare the model predictions. According to values in (Table 3), the predictions...
for all three models approach the reported infected data 2016–2018 very well. Hence, it can be said that the model solution is reliable to predict the future behaviour of TB. The error percentages between 2005 and 2018 have been calculated as 0.83%, 0.83%, and 0.45% for the SIR, the SEIR, and the BSEIR models, respectively, with the help of the $L_2$ norm. As indicated above, the obtained results show that the BSEIR model is more accurate than the SIR and the SEIR model.

On the other hand, the epidemic models which are discussed and our methodology have some limitations. The main limitations are to not assuming age-structure, emigration, seasonality, all individuals in the population have the same immune response and the same reaction to the disease for all three models. In addition to these drawbacks, the SIR model does not take into account the special features introduced by the presence of a large set of exposed individuals and, therefore, in the case of too much-exposed individuals, this makes poor predictions. Looking at the limitations of our methodology, the main limitation for all three models is to have limited data for predicting. We used the reported infected data from 2005 to 2015 since data for 2004 and before were not properly listed according to years. Besides, some initial conditions were assumed according to the examples in the literature. This uncertainty has affected the prediction results directly. Therefore, to observe the errors of the predictions caused by the assumed parameters, the sensitivity analysis has been performed. In addition to these disadvantages, fitting the parameters to the reported data has required a bit of computational work.

The dynamical behaviour of the model is defined by the basic reproduction number, $R_0$, which is the average number of new infectious caused by a single infective. Since $R_0$ depends on the model parameters, to investigate the effect of the parameters on disease transmission, the parameters $p$, $k$, $\beta$ and $y$ were subjected to sensitivity analysis. (Figures 6–8)) show the effect of parameter variation on the disease for the SIR, the SEIR and the BSEIR models, respectively. Each figure displays the number of infected individuals with respect to parameter values in (Table 1) and the corresponding curves with a specific parameter increase of 2%. Clearly from (Figures 6–8) the decrease in the total number of patients is possible by increasing the recovery rate $y$ or decreasing the transmission parameter $\beta$. Considering the BSEIR model in Fig. 8, a certain increase in the number of patients is observed when the rate of successful vaccination $p$ decreases. At the same time, the number of patients decreases when the parameter $k$, which refers to the rate of loss of effect of the vaccine, decreases. Besides, sensitivity indicates have been given in (Table 2). According to values in (Table 2), the parameter has a positive sign which means that $R_0$ increases with the parameter. While the

| Parameter | Sensitivity index $^a$ | Sensitivity index $^b$ | Sensitivity index $^c$ |
|-----------|-----------------------|-----------------------|-----------------------|
| $p$       | -                     | -                     | -0.0792               |
| $k$       | -                     | -                     | +0.0792               |
| $\beta$   | +1                    | +1                    | +1                    |
| $\mu$     | -0.0062               | -0.0086               | -0.0087               |
| $\varepsilon$ | -                 | +0.0034               | +0.0034               |
| $y$       | -0.9946               | -0.7989               | -0.9640               |

$^a$SIR Model  
$^b$SEIR Model  
$^c$BSEIR Model
parameter has a negative sign means that $R_0$ decreases for higher values of the parameter. Further, this implies that decreasing (or increasing) of the parameter $\beta$ by 10% will decrease (or increase) the basic reproductive number by 10%. Also, increasing parameter $\gamma$ by 10% will decrease the basic reproduction number by 9.946%.

Finally, a statistical evaluation of the confidence of the predictions of the SIR, the SEIR, and the BSEIR models given by the %95 confidence interval estimate has been plotted in (Figure 9).

\textbf{Figure 8.} The total number of infected individuals for different values of $p$, $k$, $\beta$ and $\gamma$ (BSEIR model).
Figure 9. Confidence intervals for the SIR, the SEIR and the BSEIR models.
4. Conclusion

In this research, the transmission dynamics of TB in Turkey have been analysed and discussed. For this purpose, three epidemic models (the SIR, the SEIR, and the BSEIR) have investigated by describing the spread of the epidemic in a certain population. By presenting the modified SIR model in which the population is fixed and the death and birth rates are equal has been started. After that, the SEIR model which includes the exposed group has been discussed. Finally, a more comprehensive model included the vaccination rate, the BSEIR, has been investigated and compared with the other models. The results showed that the obtained predictions by the BSEIR model more accurate than the other models. Since BCG vaccination is used in Turkey for a long time, the BSEIR model reveals more realistic predictions for the dynamics of the disease. The model parameters have been determined through the least square method by fitting the reported infected data. Besides, the mathematical analysis for all three models in terms of stability analysis and the importance of the reproduction number \( R_0 \) have been discussed. As indicated before, the basic reproduction number has a key role in classifying the dynamical behaviour of the models. According to calculations, \( R_0 < 1 \) has been calculated for all three models. It means that the status of TB in Turkey does not lead to an epidemic. However, if necessary precautions do not take, the infected people may increase, because the disease has not eradicated yet. Besides, the sensitivity analysis of the model parameters is performed and discussed in detail. Finally, a statistical interpretation has been added by the confidence interval estimate. It is believed that this study helps to describe the course of TB disease and determine the precautions strategy.

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ORCID

Yasin Ucakan http://orcid.org/0000-0001-5743-1862
Seda Gulen http://orcid.org/0000-0001-7092-0628
Kevser Koklu http://orcid.org/0000-0002-8609-8787

References

[1] D. Bernoulli. Essai D’une Nouvelle Analyse De La Mortalité Causée Por La Petite Vérole Et Des Avantages De L’inoculation Pour La Prévenir. In: Mem. Math. Phys. Acad. Roy. Sci. Paris. (1760). 1–45.
[2] W.H. Hamer, Epidemic disease in England, Lancet 1 (1906), pp. 733–739.
[3] R. Ross, The Prevention of Malaria, 2nd ed., Murray, London, 1911.
[4] R. Ross, An application of the theory of probabilities to the study of a priori pathometry: I, Proc. R. Soc. London A 92 (1916), pp. 204–230.
[5] R. Ross and H.P. Hudson, An application of the theory of probabilities to the study of a priori pathometry: II, Proc. R. Soc. London A 93 (1917), pp. 212–224.
[6] R. Ross and H.P. Hudson, An application of the theory of probabilities to the study of a priori pathometry: III, Proc. R. Soc. London A 93 (1917), pp. 225–240.
[7] E. Martini, Berechnungen Und Beobachtungen Zur Epidemiologie Und Bekampfung Der Malaria, W. Gente, Hamburg, 1921.
[8] A.J. Lotka, Martini’s equation for the epidemiology of immunizing diseases, Nature 111 (2793) (1923), pp. 633–634. doi:10.1038/111633a0.
[9] A.J. Lotka, Elements of physical biology, Williams& Wilkens, Baltimore (1925). Reprinted asElements of Mathematical Biology, Dower, New York, 1956.
[10] W.O. Kermack and A.G. McKendrick, A contribution to the mathematical theory of epidemics, Proc, R. Soc. Lond. A 115 (1927), pp. 700–721.
[11] M.A.B. Deakin, A standard form for the Kermack-McKendrick epidemic equation, Bull. Math. Biol. 37 (1975), pp. 91–95. doi:10.1016/S0025-5564(75)80011-5.
[12] H.W. Hethcote, Qualitative analyses of communicable disease models, Math. Biosci 28 (3–4) (1976), pp. 335–356. doi:10.1016/0025-5564(76)90132-2.
[13] A.D. Liddo, A S-I-R vector disease model with delay, Mathematical Modelling 7 (5–8) (1986), pp. 793–802. doi:10.1016/0270-0255(86)90135-1.
[14] V. Capasso and G. Serio, A generalization of the Kermack-McKendrick deterministic epidemic model, Math. Biosci 42 (1–2) (1978), pp. 43–61. doi:10.1016/0025-5564(78)90006-8.
[15] L.J. Allen, M.A. Jones, and C.F. Martin, A discrete-time model with vaccination for a measles epidemic, Math. Biosci 105 (1) (1991), pp. 111–131. doi:10.1016/0025-5564(91)90051-J.
[16] M. El-Doma, Stability analysis of a general age-dependent vaccination model of a vertically transmitted disease, International Centre for Theoretical Physics. Trieste (Italy), 1995.
[17] B. Shulgin, L. Stone, and Z. Agur, Pulse vaccination strategy in the SIR epidemic model, Bull. Math. Biol. 60 (6) (1998), pp. 1123–1148. doi:10.1016/S0092-8240(98)90005-2.
[18] A. d’Onofrio, Pulse vaccination strategy in the sir epidemic model: Global asymptotic stable eradication in presence of vaccine failures, Math. Comput. Model. 36 (4–5) (2002), pp. 473–489. doi:10.1016/S0895-7177(02)00177-2.
[19] L. Stone, B. Shulgin, and Z. Agur, Theoretical examination of the pulse vaccination policy in the SIR epidemic model, Math. Comput. Model. 31 (4–5) (2000), pp. 207–215. doi:10.1016/S0895-7177(00)00040-6.
[20] J. Satsuma, R. Willox, A. Ramani, B. Grammaticos, and A.S. Carstea, Extending the SIR epidemic model, Phys. A 336 (3–4) (2004), pp. 369–375. doi:10.1016/j.physa.2003.12.035.
[21] Y.N. Kryrchko and K.B. Bryuss, Global properties of a delayed SIR model with temporary immunity and nonlinear incidence rate, Nonlinear Anal Real World Appl 6 (3) (2005), pp. 495–507. doi:10.1016/j.nonrwa.2004.10.001.
[22] B. Buonomo, A. d’Onofrio, and D. Lacitignola, Global stability of an SIR epidemic model with information dependent vaccination, Math. Biosci 216 (1) (2008), pp. 9–16. doi:10.1016/j.mbs.2008.07.011.
[23] S. Side, A susceptible-infected-recovered model and simulation for transmission of tuberculosis, Adv Sci Lett 21 (2) (2015), pp. 137–139. doi:10.1166/asl.2015.5840.
[24] A. Takahashi, J. Spreadbury, and J. Scotti, Modeling the spread of tuberculosis in a closed population, Laporan Hasil Penelitian, 2010.
[25] D. Greenhalgh, Some results for an SEIR epidemic model with density dependence in the death rate, IMA J Math Appl Med Biol 9 (2) (1992), pp. 67–106. doi:10.1093/imammb/9.2.67.
[26] Y.A. Kuznetsov and C. Piccardi, Bifurcation analysis of periodic SEIR and SIR epidemic models, J. Math. Biol 32 (2) (1994), pp. 109–121. doi:10.1007/BF00163027.
[27] M.Y. Li and J.S. Muldowney, Global stability for the SEIR model in epidemiology, Math. Biosci 125 (2) (1995), pp. 155–164. doi:10.1016/0025-5564(95)92756-5.
[28] B. Bolker and B. Grenfell, *Space, persistence and dynamics of measles epidemic*, Philos. Trans. R. Soc. Lond., B, Biol. Sci. 348 (1995), pp. 309–320.
[29] L. Esteva and C. Vargas, *Analysis of a dengue disease transmission*, Math. Biosci 150 (2) (1998), pp. 131–151. doi:10.1016/S0025-5564(98)10003-2.
[30] J.L. Aron and I.B. Schwartz, *Seasonality and period-doubling bifurcations in an epidemic model*, J. Theor. Biol. 110 (4) (1984), pp. 665–679. doi:10.1016/S0022-5193(84)80150-2.
[31] M.Y. Li, J.R. Graef, L. Wang, and J. Karsai, *Global dynamics of a SEIR model with varying total population size*, Math. Biosci. 160 (2) (1999), pp. 191–213. doi:10.1016/S0025-5564(99)00030-9.
[32] E.A. Newton and P.A. Reiter, *Model of transmission of dengue fever with an evaluation of the impact of ultra-low volume (ULV) insecticide applications on dengue epidemics*, American Journal of Tropical Medicine and Hygiene 47 (6) (1992), pp. 709–720. doi:10.4269/ajtmh.1992.47.709.
[33] C. Castillo-Chavez and Z. Feng, *Mathematical models for the disease dynamics of tuberculosis*, In Proceedings of Fourth International Conference on Mathematical Population Dynamics, December 1995. Castillo-Chavez, C., Feng, Z, *Mathematical models for the disease dynamics of tuberculosis*, in Advances in Mathematical Population Dynamics - Molecules, Cells and Man, O. Arino, D. Axelrod, M. Kimmel, eds., Ser. Math. Biol. Med., Vol.6, World Scientific Publishing, Singapore, New Jersey, London, Hong Kong, 1997; pp. 629–656.
[34] G. Röst and J. Wu, *SEIR epidemiological model with varying infectivity and infinite delay*, Math. Biosci Eng 5 (2008), pp. 389–402.
[35] I.K. Dontwi, W. Obeng-Denteh, E.A. Andam, and L. Obiri-Apraku, *A mathematical model to predict the prevalence and transmission dynamics of tuberculosis in Amensia west district, Ghana*, British Journal of Mathematics and Computer Science 4 (3) (2014), pp. 402–425. doi:10.9734/BJMCS/2014/4681.
[36] Y. Yang, S. Tang, X. Ren, H. Zhao, and C. Guo, *Global stability and optimal control for a tuberculosis model with vaccination and treatment*, Discrete Contin. Dyn. Syst. Ser. B 21 (3) (2016), pp. 1009–1022.
[37] S. Side, S. Wahidah, M.K. Aidid, and S. Sidjara, *Global stability of SIR and SEIR model for tuberculosis disease transmission with lyapunov function method*, Asian Journal of Applied Sciences 9 (3) (2016), pp. 87–96. doi:10.3923/ajaps.2016.87.96.
[38] J. Zhang, Y. Li, and X. Zhang, *Mathematical modeling of tuberculosis data of China*, J. Theor. Biol. 365 (2015), pp. 159–163. doi:10.1016/j.jtbi.2014.10.019.
[39] C. Xu, X. Wei, J. Ciu, and X. Wang, *Mixing in regional-structure model about the influence of floating population and optimal control about TB in guangdong province of China*, Int. J. Biomath 10 (8) (2017), pp. 1750106. doi:10.1142/S1793524517501066.
[40] C.C. McCluskey, *Global stability for an SEIR epidemiological model with varying infectivity and infinite delay*, Math. Biosci Eng 6 (2009), pp. 603–610.
[41] D. Gao and N. Huang, *A note on global stability for a tuberculosis model*, Appl. Math. Lett. 73 (2017), pp. 163–168. doi:10.1016/j.aml.2017.05.004.
[42] Y. Yang, S. Tang, X. Ren, H. Zhao, and C. Guo, *Global stability and optimal control for a tuberculosis model with vaccination and treatment*, Discrete Contin. Dyn. Syst. Ser. B 21 (3) (2016), pp. 1009–1022.
[43] Y. Yang, J. Wu, J. Li, and X. Xu, *Tuberculosis with relapse: A model*, Math. Popul. Stud 24 (1) (2017), pp. 3–20. doi:10.1080/08898463.2014.998550.
[44] M. Bannon and A. Finn, *BCG and tuberculosis*, Arch. Dis. Child. 80 (1) (1999), pp. 80–83. doi:10.1136/adc.80.1.180.
[45] H. McShane, *Vaccine strategies against tuberculosis*, Swiss Med Wkly 139 (11–12) (2009), pp. 156–160.
[46] C. Castillo-Chavez and Z. Feng, *Global stability of an age-structure model for tb and its applications to optimal vaccination strategies*, Math Biosci 151 (2) (1998), pp. 135–154. doi:10.1016/S0025-5564(98)10016-0.
[47] C.H. Xiong, X.F. Liang, and H.Q. Wang, A systematic review on the protective efficacy of BCG against children tuberculosis meningitis and miliary tuberculosis, Zhongguo Ji Hua Mian Yi 15 (2009), pp. 358–362.
[48] S. Yang, X. Tang, H. Ren, and C. Guo, Global stability and optimal control for a tuberculosis model with vaccination and treatment, Discrete Contin. Dyn. Syst. Ser. B 21 (3) (2016), pp. 1009–1022. doi:10.3934/dcdsb.2016.21.1009.
[49] X. Tian, R. Xu, and J. Lin, Mathematical analysis of a cholera infection model with vaccination strategy, Appl. Math. Comput. 361 (2019), pp. 517–535.
[50] S. Liu, Y. Bi, and Y. Liu, Modeling and dynamic analysis of tuberculosis in mainland China from 1998 to 2017: The effect of DOTS strategy and further control, Theor Biol Med Model. 17 (6) (2020). doi:10.1186/s12976-020-00124-9
[51] S. Liu, Y. Bi, and Q. Huang, Mixed vaccination strategy for the control of tuberculosis: A case of study in China, Math. Biosci Eng 14 (2017), pp. 695–708.
[52] S.A. Egbetade and M.O. Ibrahim, Modelling the impact of BCG vaccines on tuberculosis epidemic, Journal of Mathematical Modelling and Application 1 (2014), pp. 49–55.
[53] Y.M. Rangkuti, M.S. Sinaga, F. Marpaung, and S. Side, A VSEIR model for transmission of TB disease in North Sumatera, Indonesia, AIP Conference Proceedeings. Langkawi, Kedah Malaysia, December 2014. DOI: 10.1063/1.4903584.
[54] A.O. Egonmwan and D. Okuonghae, Mathematical analysis of a tuberculosis model with imperfect vaccine, Int. J. Biomath 12 (7) (2019), pp. 1950073. doi:10.1142/S1793524519500736.
[55] K. Ergen, A. Cilli, and N. Yahnioglu, Predicting epidemic diseases using mathematical modelling of SIR, Acta Phys. Polon. A 128 (2B) (2015), pp. 273–275. doi:10.12693/ APhysPolA.128.B-273.
[56] A. Cilli and K. Ergen, Investigation of various infectious diseases in Turkey by mathematical models SI and SIS, International Journal of Computational and Experimental Science and Engineering 5 (2) (2019), pp. 72–75. doi:10.22399/ijcesen.542383.
[57] WHO, Global tuberculosis report 2007, World Health Organization (2007). Available at https://apps.who.int/iris/bitstream/handle/10665/43629/9789241563141_eng.pdf?sequence=1 .
[58] WHO, Global tuberculosis report 2008, World Health Organization (2008). Available at https://apps.who.int/iris/bitstream/handle/10665/43831/9789241563543_eng.pdf?sequence=1 .
[59] WHO, Global tuberculosis report 2009, World Health Organization (2009). Available at https://apps.who.int/iris/bitstream/handle/10665/44035/9789241563802_eng.pdf?sequence=1 .
[60] WHO, Global tuberculosis report 2010, World Health Organization (2010). Available at https://apps.who.int/iris/bitstream/handle/10665/44425/9789241564069_eng.pdf?sequence=1 .
[61] WHO, Global tuberculosis report 2011, World Health Organization (2011). Available at https://apps.who.int/iris/bitstream/handle/10665/44728/9789241564380_eng.pdf?sequence=1 .
[62] WHO, Global tuberculosis report 2012, World Health Organization (2012). Available at https://apps.who.int/iris/bitstream/handle/10665/75938/9789241564502_eng.pdf?sequence=1 .
[63] WHO, Global tuberculosis report 2013, World Health Organization (2013). Available at https://apps.who.int/iris/bitstream/handle/10665/91355/9789241564656_eng.pdf?sequence=1 .
[64] WHO, Global tuberculosis report 2014, World Health Organization (2014). Available at https://apps.who.int/iris/bitstream/handle/10665/137094/9789241564809_eng.pdf?sequence=1 .
[65] WHO, Global tuberculosis report 2015, World Health Organization (2015). Available at https://apps.who.int/iris/bitstream/handle/10665/191102/9789241565059_eng.pdf?sequence=1 .
[66] WHO, *Global tuberculosis report 2016*, World Health Organization (2016). Available at https://apps.who.int/iris/bitstream/handle/10665/250441/9789241565394eng.pdf?sequence=1.

[67] WHO, *Global tuberculosis report 2017*, World Health Organization (2017). Available at http://apps.who.int/iris/bitstream/10665/259366/1/9789241565516eng.pdf?ua=1.

[68] L.J.S. Allen, *An Introduction to Mathematical Biology*, NJ Pearson, Upper Sandle River, 2007.

[69] O. Bjørnstad, *SEIR model*, McMaster University 2005. p. 1–8. Available at https://ms.mcmaster.ca/bolker/eedis/sir.pdf.

[70] F.R. Gantmacher. *Matrix Theory*. Interscience.Vol. II. Chelsea Pub. Co.: New York. 1964.

[71] P.V.D. Driessche and J. Watmough, *Reproduction numbers and sub-threshold endemic equilibria for compartmental models of disease transmission*, Math. Biosci 180 (1–2) (2002), pp. 29–48. doi:10.1016/S0025-5564(02)00108-6.

[72] *Tuberculosis in Turkey 2017 report*, Republic of Turkey Ministry of Health (2017). Available at https://hsgm.saglik.gov.tr/depo/haberler/verem-savas-raporu-2016-2017/Turkiyede_Verem_Savasi_2017_Raporu.pdf

[73] *The ministry of health of Turkey health statistics yearbook*, Refik Saydam Hygiene Center Presidency School of Public Health (2010). Available at http://sbu.saglik.gov.tr/ekutuphane/kitaplar/saglikistatistikleriyyilligi2010.pdf.

[74] Turkish Statistical Institute. Available at https://tuikweb.tuik.gov.tr/PreIstatistikTablo.do?istab_id=1592.

[75] S. Whang, S. Choi, and E. Jung. *A dynamic model for tuberculosis transmission and optimal treatment strategies in South Korea*, J. Theoret. Biol 279 (1) (2011), pp. 120–131. doi:10.1016/j.jtbi.2011.03.009.