An Intrinsic Analysis of Human Brucellosis Dynamics in Africa

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Author’s contributions

This work was carried out in collaboration between both authors. Both authors read and approved the final manuscript.

Abstract

Brucellosis is one of the most common zoonotic infections globally. It affects humans, domestic animals and wildlife. In this paper, we conduct an intrinsic analysis of human brucellosis dynamics in non-periodic and periodic environments. As such we propose and study two mathematical models for human brucellosis transmission and control, in which humans acquire infection from cattle and wildlife. The first model is an autonomous dynamical system and the second is a non-autonomous dynamical system in which the seasonal transmission of brucellosis is incorporated. Disease intervention strategies incorporated in this study are cattle vaccination, culling of infectious cattle and human treatment. For both models we conduct both epidemic and endemic analysis, with a focus on the threshold dynamics characterized by the basic reproduction numbers. Using sensitivity analysis we established that $R_0$ is most sensitive to the rate of brucellosis transmission from buffalos to cattle, the result suggest that in order to control human brucellosis there is a need to control cattle infection. Based on our models, we also formulate an optimal control problem with cattle vaccination and culling of infectious cattle as control functions. Using reasonable parameter values, numerical simulations of the optimal control demonstrate the possibility of reducing brucellosis incidence in humans, wildlife and cattle, within a finite time horizon, for both periodic and non-periodic environments.

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1 Introduction

Globally, human brucellosis remains an important and widespread infection [1]. The infection is more common in Mediterranean areas, the south and the center of America, Africa, Asia, Arab peninsula, Indian subcontinent and the Middle East [2]. In 2012, reported brucellosis incidence in some endemic regions were as follows: Saudi Arabia (214.4), Iran (238.6), Turkey (262.2), Iraq (278.4), and Syria (1603.4) [3]. However, the World Health Organization (WHO) believes that the real incidence is 10–25 times more than what have been reported [2]. Although, human brucellosis in rampant in many developing nations it is also a severe public health problem in China where 160, 214 brucellosis cases were observed in the period 2005–2010 [4].

In animals, brucellosis is transmitted by direct contact transmission through the brucella carriers or indirect contact transmission when animals ingest contaminated forages or the excrement containing large quantities of bacteria, generally discharged by infected animals [5]. Domesticated species such as cattle, sheep, horse and goats are regarded as the main source of human brucellosis [2], in which transmission may occur directly or through the consumption of unpasteurised dairy products [5]. The cross-transmission of brucellosis between domesticated animals and wildlife is well documented [6, 7]. However, the debate on whether wildlife is the reservoir of infection for domestic animals or vice versa continues [6]. One wild animal that is a villain for inter-species spread of many infectious disease such as brucellosis, foot-and-mouth disease-virus (FMDV) in many Africa nations is the African buffalo [6]. African buffaloes have several intrinsic behavioural characteristics which are key to inter-species spread of infectious diseases. They are highly mobile and sociable species and they often move in large herds of 1000 or more [8].

Recently, a number of veterinary scientists have suggested that buffalo, a preferred source of bush meat could be another source of human brucellosis in many developing nations [6]. Buffalo meat is highly preferred bush meat in many African countries [9]. Since bush meat is consumed and handled (legally and illegally) in many developing nations its contribution to human brucellosis cannot be ignored.

Several mathematical models have been proposed to study the dynamics of brucellosis outbreaks [5, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26]. Undeniably, these studies have produced many useful results and improved the existing knowledge on brucellosis dynamics. One of the limitations of these models, however, is that none of them incorporated the aspect of bush meat on modeling the transmission dynamics of human brucellosis. In this paper we develop a novel mathematical model that evaluates the impact of bush meat on human brucellosis dynamics. Our model will incorporate human population, cattle and wildlife (African buffalo). In addition, we will explore optimal disease control measures based on cattle vaccination, culling of infectious cattle and treatment of infected humans.

The remainder of this paper is organized as follows. In section 2 we formulate and comprehensively analyze the transmission dynamics of brucellosis in non-periodic environments. We provide sensitivity analysis of the basic reproductive number on various model parameters in non-periodic environments, identifying the parameters to which reproductive number is most sensitive, we use this information to suggest strategies for controlling brucellosis using techniques from optimal control theory. We extend the autonomous brucellosis models formulated to incorporate seasonal variations on disease transmission. We then conduct mathematical analyses, including the computation of the basic
2 Materials and Methods

2.1 An autonomous brucellosis model

We propose an autonomous dynamical system that comprise of cattle population, human population and African buffalo population. The African buffalo population constitute three compartments: susceptible \( S_b(t) \), symptomatic infectious \( I_b(t) \) and carrier state - asymptomatic persistent infection \( A_b(t) \). Thus the total population of buffaloes at time \( t \) is \( N_b(t) = S_b(t) + I_b(t) + A_b(t) \). The cattle population is subdivided into classes of: susceptible \( S_c(t) \), infectious \( I_c(t) \), and the total cattle population at time \( t \) is \( N_c(t) = S_c(t) + I_c(t) \). Further, the human population constitute: susceptible \( S_h(t) \) and infectious \( I_h(t) \). The total human population at time \( t \) is \( N_h(t) = S_h(t) + I_h(t) \). There are some assumptions for our model:

1. Brucellosis in the exposure period is hardly detected, and animals in this period can also infect susceptible animal and humans. Hence, we ignored the exposed/latent period in both human and animal population (see for example [15, 20, 21, 22]);

2. Prior studies suggest that African buffaloes have more chances of becoming chronic carriers of the disease [6, 27]. As such we omitted the carrier compartment on describing brucellosis transmission dynamics in cattle and humans;

3. Here, brucellosis transmission rate is being modeled by the mass action incidence since it is appropriate when \( N(t) \) is not too large [28]. We assume that the transmission rate is dependent on the size of the population which implies that the contact rate is an increasing function of the population. The mass action incidence is density-dependent since contact rate per infective is proportional to the density of the infectious host.

In all the discussions to follow we will denote the African buffalo, cattle and human by subscripts \( b, c \) and \( h \), respectively. A flow diagram describing the model is given in Fig.1 and the model equations are:

\[
\begin{align*}
\frac{dS_b}{dt} &= \Lambda_b - [\beta_{bh}(I_b + \epsilon A) + \beta_{hb} I_h] S_b - \mu_b S_b, \\
\frac{dI_b}{dt} &= [\beta_{bh}(I_b + \epsilon A) + \beta_{hb} I_h] S_b - [\mu_b + \gamma] I_b, \\
\frac{dA_b}{dt} &= f\gamma I_b - [\mu_b + d_b] A, \\
\frac{dS_c}{dt} &= \Lambda_c - [\beta_{ch}(I_h + \epsilon A) + \beta_{hc} I_c] S_c - [\sigma + \mu_c] S_c, \\
\frac{dI_c}{dt} &= [\beta_{ch}(I_h + \epsilon A) + \beta_{hc} I_c] S_c - [\mu_c + d_c] I_c, \\
\frac{dS_h}{dt} &= \Lambda_h - [\beta_{hc}(I_c + \epsilon A) + \beta_{ch} I_c] S_h - [\mu_h + d_h] S_h, \\
\frac{dI_h}{dt} &= [\beta_{hc}(I_c + \epsilon A) + \beta_{ch} I_c] S_h - [\mu_h + d_h + \theta] I_h, \\
\frac{dA_h}{dt} &= [\beta_{hc}(I_c + \epsilon A)S_h + \beta_{ch} I_c S_h - (\theta + \mu_h) A_h, \\
\end{align*}
\]

In system (1), \( \Lambda_b \) denote the constant recruitment rate into the population through birth, \( \mu_i \) is the natural-related death rate \((i = b,c,h)\), \( d_b \) is the disease-related mortality rate for buffalo population, \( d_c = \alpha + \delta \), where \( \alpha \) is the culling rate and \( \delta \) is the disease-related mortality rate, \( \epsilon \) is the vaccination rate, \( \epsilon \) accounts for the unequal chances of disease transmission between symptomatic and asymptomatic buffaloes. \( \beta_{ij} \) \((i, j = b,c)\) denotes disease transmission rate, with \( i = j \) implying buffalo-to-buffalo or cattle-to-cattle transmission and \( i \neq j \) signify cross-transmission, respectively. \( \beta_{bh} \) and \( \beta_{ch} \) denotes transmission rate from buffalo and cattle, respectively, to humans. Further, infected African buffaloes display clinical signs of the disease for \( \gamma^{-1} \) days after which a fraction \( f \) become chronic carriers and the complementary \((1 - f)\) succumb to disease-related death. Infected human displaying clinical signs of the disease are treated at rate \( \theta \). Prior studies suggests that the optimal treatment of uncomplicated brucellosis should be based on a six-week regimen of doxycycline.
combined either with streptomycin for 2–3 weeks, or rifampicin for six weeks [29].

Although, species can be infected by the brucella through indirect transmission (environmental transmission), prior studies suggests that this form of infection plays a relatively small role on the spread of brucellosis [11, 14]. In addition, prior studies also suggests that humans rarely transmit the disease [11, 14].

![Flow diagram representing the transmission routes and other processes model by system (1).](image)

It can easily be verified that model (2) has a unique and bounded solutions with initial value in $\mathbb{R}_5^+$. Further, the compact set

$$\Gamma = \left\{ (S_b, I_b, A, S_c, I_c) \in \mathbb{R}_5^+ : N_b \leq \frac{\Lambda_b}{\mu_b}, N_c \leq \frac{\Lambda_c}{\mu_c} \right\},$$

is positively invariant and attracting with respect to model (2).

### 2.1.1 The reproductive number

It is evident that (2) always has a disease-free equilibrium (DFE) given by

$$\mathcal{E}^0 : [S_b^0, I_b^0, A^0, S_c^0, I_c^0] = \left[ \frac{\Lambda_b}{\mu_b}, 0, 0, \frac{\Lambda_c}{\sigma + \mu_c} \right].$$

One measure of the severity of a disease is the basic reproductive number, $\mathcal{R}_0$, which is defined as the average number of secondary infections caused by a single infected animal in a completely susceptible population. Using the second generation matrix approach [30], the non-negative matrix
Thus, the next generation matrix of system (2) is

\[
F = \begin{bmatrix}
\frac{\beta_{ab}\Lambda_b}{\mu_b} & \frac{\beta_{ab}\Lambda_b}{\mu_b} & \frac{\beta_{ab}\Lambda_b}{\mu_b} \\
\frac{\beta_{bc}\Lambda_c}{\sigma+\mu_c} & \frac{\beta_{bc}\Lambda_c}{\sigma+\mu_c} & \frac{\beta_{bc}\Lambda_c}{\sigma+\mu_c} \\
\frac{\beta_{cc}\Lambda_c}{\mu_c} & \frac{\beta_{cc}\Lambda_c}{\mu_c} & \frac{\beta_{cc}\Lambda_c}{\mu_c}
\end{bmatrix}, \quad \text{and} \quad V = \begin{bmatrix}
(\mu_b + \gamma) & 0 & 0 \\
-f\gamma & (\mu_b + d_b) & 0 \\
0 & 0 & (\mu_c + d_c)
\end{bmatrix}.
\]

Thus, the next generation matrix of system (2) is

\[
FV^{-1} = \begin{bmatrix}
M_{11} & M_{12} & M_{13} \\
M_{21} & M_{22} & M_{23} \\
0 & 0 & 0
\end{bmatrix},
\]

where

\[
\begin{align*}
M_{11} &= \frac{\Lambda_b\beta_{ab}}{(\mu_b + \gamma)\mu_b} + \frac{f\gamma\Lambda_c\beta_{bc}}{(\mu_b + \gamma)(\mu_b + d_b)\mu_b}, \\
M_{21} &= \frac{\Lambda_c\beta_{bc}}{(\mu_b + \gamma)(\sigma + \mu_c)} + \frac{f\gamma\Lambda_c\beta_{bc}}{(\mu_b + \gamma)(\mu_b + d_b)(\sigma + \mu_c)}, \\
M_{12} &= \frac{\Lambda_c\beta_{bc}}{\mu_b(\mu_b + d_b)}, \quad M_{22} = \frac{\Lambda_c\beta_{bc}}{(\sigma + \mu_c)(\mu_b + d_b)}, \\
M_{13} &= \frac{\Lambda_b\beta_{ab}}{\mu_b(\mu_b + d_b)}, \quad M_{23} = \frac{\Lambda_c\beta_{bc}}{(\sigma + \mu_c)(\mu_b + d_b)}.
\end{align*}
\]

It follows that the basic reproductive number is

\[
R_0 = \rho(FV^{-1}) = \frac{M_{11} + M_{22} + \sqrt{(M_{11} - M_{22})^2 + 4M_{12}M_{23}}}{2},
\]

Defining the appropriate value of \(R_0\) for a disease characterized by hidden infections is challenging but essential in developing control measures. The reproductive number is the key threshold parameter whose values determine the global dynamics of system (2). A disease is considered to be endemic if \(R_0 > 1\). However, if \(R_0 \leq 1\) it implies that the disease dies out.

### 2.1.2 Equilibrium analysis

**Theorem 2.1.** If \(R_0 \leq 1\), the system (2) has a unique DFE that is globally asymptotically stable in the region \(\Gamma\).

**Proof.** Let \(y(t) = [I_b(t), A(t), I_c(t)]\). Since

\[
\begin{align*}
\dot{I}_b &\leq [\beta_{ab}(I_b + \epsilon A) + \beta_{bc}I_c]S_b - [\mu_b + \gamma]I_b, \\
\dot{A} &\leq f\gamma I_b - [\mu_b + d_b]A, \\
\dot{I}_c &\leq [\beta_{bc}(I_b + \epsilon A) + \beta_{cc}I_c]S_c - [\mu_c + d_c]I_c,
\end{align*}
\]

it follows that

\[
\dot{y} \leq (F - V)y,
\]

where \(F\) and \(V\) are defined in Eq. (4). One can easily deduce that, both \(F\) and \(V^{-1}\) are non-negative. By the Perron-Frobenius Theorem, the non-negative matrix \(V^{-1}F\) has a non-negative
It can easily be verified that model (6) has a unique endemic equilibrium \( \mathcal{E}^* \), and that the global stability of \( \mathcal{E}^* \) is the same as that of system (2). Consider the Lyapunov function

\[
\mathcal{L} = S_b^* [x - 1 - \ln x] + I_b^* [y - 1 - \ln y] + \frac{[\beta_{bb}A^*S_b^* + \beta_{bc}A^*S_c^*]A^*}{f\gamma I_b^*} [z - 1 - \ln z] + S_c^* [u - 1 - \ln u] + I_c^* [v - 1 - \ln v].
\]
In this section we perform the sensitivity analysis of the model system (1). The threshold quantity $R_0$ is an important parameter to determine the persistence and extinction of brucellosis disease transmission in the population. To be able to suggest the most efficient way of controlling the disease we need to determine the parameters we can control and to

Differentiating $\mathfrak{U}$ with respect to $t$ along solutions of (6) gives:

$$\frac{d\mathfrak{U}}{dt} = (x - 1) \left[ \lambda_s \left( \frac{1}{x} - 1 \right) - \beta_{bc} I_b^* S_b^*(y - 1) - \beta_{bc} \epsilon A^* S_b^*(z - 1) - \beta_{cb} I_c^* S_b^*(v - 1) \right]$$

$$+ (y - 1) \left[ \beta_{bc} I_b^* S_b^*(x - 1) + \beta_{bc} \epsilon A^* S_b^* \left( \frac{2x}{y} - 1 \right) + \beta_{cb} I_c^* S_b^* \left( \frac{v}{y} - 1 \right) \right]$$

$$+ (u - 1) \left[ u \left( \frac{1}{u} - 1 \right) - \beta_{bc} I_b^* S_b^*(y - 1) - \beta_{bc} \epsilon A^* S_b^*(z - 1) - \beta_{cb} I_c^* S_b^*(v - 1) \right]$$

$$+ (v - 1) \left[ \beta_{bc} I_b^* S_b^*(u - 1) + \beta_{bc} \epsilon A^* S_b^* \left( \frac{2u}{v} - 1 \right) + \beta_{cb} I_c^* S_b^* \left( \frac{yu}{v} - 1 \right) \right]$$

$$+ \left[ \beta_{bc} \epsilon A^* S_b^* + \beta_{bc} \epsilon A^* S_b^* \right] \left( \frac{y}{x} - 1 \right)$$

$$= F(x, y, z, u, v).$$

At endemic point we have the following identities:

$$\mu_b = \frac{\lambda_b}{S_b} - \beta_{bc} I_b^* S_b^* + \beta_{bc} \epsilon A^* S_b^*$$

$$\mu_b + \gamma = \beta_{bc} S_b^* + \beta_{bc} \epsilon A^* S_b^*$$

$$\sigma + \mu_c = \frac{\lambda_c}{S_c} - \beta_{bc} I_b^* S_b^* + \beta_{bc} \epsilon A^* S_b^*$$

$$\mu_c + d_c = \beta_{bc} S_c^* + \beta_{bc} \epsilon A^* S_b^*$$

$$\mu_b + d_b = \frac{f \gamma I_b^*}{A^*}.$$

To assure that $F(x, y, z, u, v) \leq 0$ for $x > 0, y > 0, z > 0, u > 0, v > 0$, the following condition must be satisfied $\beta_{bc} I_b^* S_b^* = \beta_{bc} (I_b^* + A^*) S_b^*$ (see [34]). After some algebraic manipulations, we have

$$\frac{d\mathfrak{U}}{dt} = (\mu_b S_b^* + \beta_{bc} I_b^* S_b^*) \left( 2 - x - \frac{1}{x} \right) + \left( \sigma + \mu_c \right) S_c^* + \beta_{bc} I_c^* S_c^* \left( 2 - u - \frac{1}{u} \right)$$

$$+ \beta_{bc} \epsilon A^* S_b^* \left( 3 - \frac{1}{x} - \frac{y}{z} - \frac{zx}{y} \right) + \beta_{bc} \epsilon A^* S_c^* \left( 4 - \frac{1}{x} - \frac{1}{u} - \frac{vx}{y} - \frac{yv}{v} \right)$$

$$+ \beta_{bc} \epsilon A^* S_c^* \left( 5 - \frac{1}{x} - \frac{1}{u} - \frac{vx}{y} - \frac{zv}{v} - \frac{zx}{z} \right).$$

Since the arithmetic mean is greater or equal to the geometric mean, it can easily be verified that $\mathfrak{U} \leq 0$ provided that $S_b^*, I_b^*, A^*, S_c^*, I_c^*$ are positive, where the equality $\mathfrak{U} = 0$ holds only for $x = y = z = u = v = 1$. Therefore $\mathfrak{U} \leq 0$ holds. Then the endemic equilibrium point $\Omega^*$ is globally asymptotically stable if $R_0 > 1$ by LaSalle’s invariance principle [32].

### 2.1.3 Sensitivity analysis of the reproduction number

In this section we perform the sensitivity analysis of the model system (1). The threshold quantity $R_0$ known as basic reproduction number is an important parameter to determine the persistence and extinction of brucellosis disease transmission in the population. To be able to suggest the most efficient way of controlling the disease we need to determine the parameters we can control and to
which \( R_0 \) is more sensitive. Therefore we perform the sensitivity analysis of the model system (1) using partial rank correlated coefficient (PRCC) developed in [33] and the values of the parameters used in the model simulations are in Table (2) to demonstrate the influence of each parameter in the size of threshold quantity \( R_0 \). PRCC is an efficient sensitivity analysis method based on sampling. PRCC assigns a value between \(-1\) to \(+1\) for each parameter. Positive PRCC value indicates a positive correlation of the parameter with the disease maintenance, whereas a negative value indicates a negative correlation with the infectiousness of the diseases. The Parameters studied are: \( \Lambda_b, \Lambda_c, \Lambda_h, \mu_b, \mu_c, \mu_h, d_b, \beta_{bc}, \beta_{bb}, \beta_{cc}, \beta_{bh}, \beta_{ch}, f, \theta, \gamma, \sigma, \beta_{hc}, \epsilon \).

**Definition 2.1.** (See, [33]) The normalized sensitivity index of \( R_0 \) which depends on differentiability of parameter, \( \omega \) is defined as follows:

\[
\Psi R_0^\omega = \frac{\partial R_0}{\partial \omega} \times \frac{\omega}{R_0}.
\]

From (9), the value of normalized sensitivity index for each parameter used in the model (1) is summarized in Table 1:

| Parameter | \( \Lambda_b \) | \( \Lambda_c \) | \( \Lambda_h \) | \( \mu_b \) | \( \mu_c \) | \( \mu_h \) | \( d_b \) |
|-----------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Index     | +0.7399         | +0.2601         | 0               | -1.0476         | -0.1141         | 0               | -0.4615         |

| Parameter | \( \beta_{bc} \) | \( \beta_{bb} \) | \( \beta_{cc} \) | \( \beta_{bh} \) | \( \beta_{ch} \) | \( f \) | \( \theta \) |
|-----------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Index     | +0.1937         | +0.7309         | 0               | 0               | 0               | +0.64           | 0               |

| Parameter | \( \gamma \) | \( \sigma \) | \( \beta_{hc} \) | \( d_c \) | \( \epsilon \) |
|-----------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Index     | -0.2308         | -0.1460         | +0.2601         | 0               | +0.7691         |

**Fig. 2.** Sensitivity analysis of the model system (1)
In Fig. 2, we observed that model parameters such as $\Lambda_b$, $\Lambda_c$, $\beta_{bc}$, $\beta_{bb}$, $\epsilon$, and $f$, have a positive influence on the $R_0$, that is, whenever they are increased, the size of $R_0$ increases. For example, an increase in recruitment rate of Buffalo $\Lambda_b$ by 73.99% will lead to an increase in the size of $R_0$ by 73.99%. In the other-hand, model parameters with negative index values have a negative influence on $R_0$, for example, an increase in mortality rate of Buffalo $d_b$ by 46.15% will lead to a decrease on the magnitude of $R_0$ by 46.15%.

![Graph](image)

(a)

![Graph](image)

(b)

Fig. 3. Effects of varying (a) progression rate of infected cattle to chronic stage modeled by parameter $\gamma$ on $R_0$ (b) vaccination rate of susceptible cattle modeled by parameter $\sigma$ on $R_0$

Numerical results in Fig. 3a shows the progression rate of infected cattle from susceptible to chronic stage modeled by parameter $\gamma$ on $R_0$. Overall, we noted that increase on progression rate of infected cattle to chronic stage reduce the size of $R_0$. In particular, one can note that whenever
the progression rate is greater than 0.5 the disease dies in the community. Fig 3b demonstrates the effect of vaccination rate of susceptible cattle on the spread of brucellosis disease in the population. Overall, we observed that whenever the vaccination rate of susceptible cattle is less than 0.5 the disease persists in the community.

Fig. 4. Effects of varying rate of modifying factor for disease transmission modeled by parameter $\epsilon$ on $R_0$

Fig. 4 demonstrates the effects of modifying factor in the dynamics of brucellosis disease transmission. Overall, one can note that whenever the modifying factor for disease transmission is less than 0.5 the magnitude of $R_0$ is less than unit and thus, the disease dies in the population.

Fig. 5. Contour plot of the basic reproduction number $R_0$ as the function of treatment rate of infected cattle (modeled by parameter $\theta$) and vaccination rate of susceptible cattle (modeled by parameter $\sigma$)
Fig. 5 shows the contour plot of basic reproduction number $R_0$ as the function of treatment rate of infected cattle (modeled by parameter $\sigma$) and vaccination rate of susceptible cattle (modeled by parameter $\nu$). Overall, we noted that, increase on vaccination rate of susceptible and treatment of infected cattle reduce the size of $R_0$. In particular one can note that whenever $\sigma$ is greater than 0.5 the disease dies in the population.

### 2.2 A periodic brucellosis model

Infectious disease dynamics are often strongly influenced by seasonal patterns, irrespective of pathogen rate transmission mode [7, 35]. The breadth and consistency of these patterns suggest that seasonal influence on host and pathogen biology can have significant effects on patterns of pathogen invasion and transmission [36]. The relationship between host abundance and pathogen transmission, influenced strongly by seasons in some environments such as the semi-arid, is central to understanding infectious disease ecology and patterns and processes of pathogen invasion [7].

In fact, like many other infectious diseases, brucellosis is significantly influenced by seasonal variations, and prior studies have demonstrated a strong connection between brucellosis infection and seasonal variations [7, 37, 38]. Factors such as the seasonal availability of forage which in turn lead to nomadic animal farming may be attributed to seasonality of brucellosis dynamics. Botswana provides an important example of this potential influence with extreme seasonal climatic variation, which occurs within and between years. In Botswana water availability is highly variable in time and space in relation to rainfall patterns and this can strongly influence density and spatial distribution of domestic animals and wildlife including buffalo over the whole year including buffalo calving periods [6].

Against this background, in this section, we extend model (1) to incorporate seasonality. Thus we introduce seasonal-induced transmission rate. Our new model takes the form:

\[
\begin{align*}
\frac{dS}{dt} &= \Lambda_S - (\beta_{bc}(t)I_b + \epsilon A) + \beta_{bc}(t)I_c)S_b(t) - \mu_S S(t), \\
\frac{dA}{dt} &= \gamma I_b - \mu_A A(t), \\
\frac{dS}{dt} &= \Lambda_S - (\beta_{bc}(t)I_b + \epsilon A) + \beta_{bc}(t)I_c)S_c(t) - (\sigma + \mu_c)S_c(t), \\
\frac{dA}{dt} &= \mu_A A(t), \\
\frac{dS}{dt} &= \Lambda_S - (\beta_{bc}(t)I_b + \epsilon A) + \beta_{bc}(t)I_c)S_c(t) - (\mu_c + \mu_d)A(t), \\
\frac{dA}{dt} &= \Lambda_S - (\beta_{bc}(t)I_b + \epsilon A)S_b - \beta_{bc}(t)I_cS_c(t) - (\mu_c + \mu_b)A(t). \\
\end{align*}
\]

All the variables and model parameters are assumed to be positive and they retain the same definitions as in model (1). Further, we assume that $\beta_{ij}(t), (i, j = b, c, h)$ are periodic continuous functions in $t$ with a period $\omega > 0$ (specifically, $\omega = 12$ months). Thus,

\[
\beta_{ij}(t) = \beta_{ij}\left[1 + a_k \sin\left(\frac{2\pi t}{12}\right)\right], \quad k = 1, 2, 3, 4, 5, 6
\]

where $\beta_{ij}$ denotes the basic contact rate without seasonal forcing and $0 < a_k < 1$ denotes the magnitude of seasonal fluctuations.

#### 2.2.1 The reproductive number

To introduce the basic reproduction number in the fluctuating environment Wang and Zhao [39], extended the general procedure presented by Driessche and Watmough [30] by introducing the next infection operator

\[
(L\phi)(t) = \int_0^\infty Y(t, t - s)F(t - s)\phi(t - s)ds
\]
Assume that $Y(t, s), t \geq s$, is the evolution operator of the linear $\omega$-periodic system $\frac{dY}{dt} = V(t)y$ and $\phi(t)$, the initial distribution of infectious animals, is $\omega$-periodic and always positive. Then effective reproductive number of the system (10) is then established by calculating the spectral radius of the next infection operator,

$$ \mathcal{R}_e = \rho(L). $$

Thus, the evolution operator $Y(t, s)$, for the system (10) is

$$ Y(t, s) = \begin{bmatrix} \frac{f_{\gamma}}{(T - d_{b})} e^{-(\mu_k + \gamma)(t-s)} - e^{-(\mu_k + \gamma)(t-s)} & 0 & 0 \\ 0 & e^{-(\mu_k + d_{b})(t-s)} & 0 \\ 0 & 0 & e^{-(\mu_k + d_{e})(t-s)} \end{bmatrix}. $$

(13)

The next infection operator can be numerically evaluated by

$$ (L\phi(t) = \int_0^\infty Y(t, t-s)F(t-s)\phi(t-s)ds = \int_0^T G(t,s)\phi(t-s)ds, $$

where

$$ G(t,s) \approx \sum_{k=0}^M Y(t, t-s - k\omega)F(t-s) $$

for some positive integer $M$ large enough, and

$$ \begin{align*}
\ell_{11} &= l_{12} = \frac{\beta_{ab}(t-s)\lambda_e}{\rho_b} e^{-(\mu_k + \gamma)(s+k\omega)}, \\
\ell_{13} &= \frac{\beta_{ak}(t-s)\lambda_b}{\rho_b} e^{-(\mu_k + \gamma)(s+k\omega)}, \\
\ell_{21} &= l_{22} = \frac{f_{\gamma}}{(T - d_{b})} \left[ \frac{\beta_{ab}(t-s)\lambda_e}{\rho_b} e^{-(\mu_k + d_{b})(s+k\omega)} - l_{11} \right], \\
l_{23} &= \left[ \frac{\beta_{ak}(t-s)\lambda_b}{\rho_b} e^{-(\mu_k + d_{b})(s+k\omega)} - l_{13} \right], \\
\ell_{31} &= l_{32} = \frac{\beta_{ae}(t-s)\lambda_e}{(\sigma + \rho_e)} e^{-(\mu_k + d_{e})(s+k\omega)}, \\
l_{33} &= \frac{\beta_{ae}(t-s)\lambda_e}{(\sigma + \rho_e)} e^{-(\mu_k + d_{e})(s+k\omega)}. 
\end{align*} $$

In the special case of $\beta_{ij}(t) \equiv \beta_{ij}, \forall t \geq 0, we obtain F(t) \equiv F_0, and V(t) \equiv V, \forall t \geq 0, then \mathcal{R}_e = \mathcal{R}_0$.

### 2.2.2 Brucellosis extinction and persistence

In this section, we present that if $\mathcal{R}_e < 1$, then DFE is globally stable and the disease dies out. Then, if $\mathcal{R}_e > 1$ the disease persist.

In the special case of $\beta_{ij}(t) \equiv \beta_{ij}, \forall t \geq 0, we obtain F(t) \equiv F_0, and V(t) \equiv V, \forall t \geq 0, then \mathcal{R}_e = \mathcal{R}_0$. It can easily be verified that system (10) satisfies assumptions (A1)-(A7) in Wang and Zhao (2008) [39]. Thus, we have the following results, which are crucial for our simulations and the main analytical results in this section.
Lemma 2.3. (Wang and Zhao Theorem 2.2 in [39]). The following statements are valid:

(i) $\mathcal{R}_0 = 1$ if and only if $\rho(\Phi_{(F-V)}(\omega)) = 1$.

(ii) $\mathcal{R}_0 > 1$ if and only if $\rho(\Phi_{(F-V)}(\omega)) > 1$.

(iii) $\mathcal{R}_0 < 1$ if and only if $\rho(\Phi_{(F-V)}(\omega)) < 1$.

We now proceed to prove Theorem 2.5.

Proof. In the case where $\mathcal{R}_s < 1$, Lemma 2.3 implies that $\mathcal{E}^0$ is locally asymptotically stable. It suffices to prove that $\mathcal{E}^0$ is globally attractive in $\Gamma$.

Assume that $\mathcal{R}_s < 1$. By Lemma 2.3, it follows that for any $\varphi > 0$, there exists large $t_0 > 0$ such that $S_0(t) < S_0^0 + \varphi$ and $S_1 < S_1^0 + \varphi$, when $t > t_0$. Then for system (10), we have, when $t > t_0$, that

\[
\begin{align*}
I_s(t) &\leq \left[\beta_s(t)(I_s + \epsilon A) + \beta_a(t)I_c\right][S_0^0(t) + \varphi] - [\mu_b + \gamma]I_s, \\
\dot{A}(t) &= f_1I_s - [\mu_b + d_2]A(t), \\
\dot{L}_c(t) &\leq \left[\beta_a(t)(I_s + \epsilon A) + \beta_c(t)L_c\right][S_0^0(t) + \varphi] - [\mu_c + d_c]L_c(t).
\end{align*}
\]

(14)

Considering the comparison system,

\[
dh(t) = (F(t) - V(t) + M_f)h(t), \quad h(t) = (I_s(t), I_c(t), A(t)).
\]

(15)

By Lemma 2.1 in [40], it follows that there exists a positive $\omega$-periodic function $\bar{h}(t)$ such that $h(t) = e^{\psi t}$ is a solution of the system (15) where $\psi = \frac{1}{M_f^*} \ln \rho(\Phi_{(F-V+M_f)}(\omega))$. Further, we know that $\mathcal{R}_s < 1$, if and only if $\rho(\Phi_{(F-V+M_f)}(\omega)) < 1$. Since $\rho(\Phi_{(F-V+M_f)}(\omega)) < 1$, it follows that, $\psi$ is a negative constant. Therefore, we have $h(t) \to 0$ as $t \to +\infty$. This implies that the zero solution of system (15) is globally asymptotically stable. For any non-negative initial value $(I_s(0), A(0), I_c(0))$ for system (14), there is a sufficient large $M^* > 0$ such that $(I_s(0), A(0), I_c(0))^T \leq M^* h(0)$ holds. Following the comparison principle [41], we have $(I_s(t), A(t), I_c(t))^T \leq M^* h(t)$ for all $t > 0$ where $M^* h(t)$ is also a solution of system (15). Therefore, we get $I_s(t) \to 0$, $A(t) \to 0$ and $I_c(t) \to 0$, as $t \to +\infty$. By the theory of asymptotic autonomous systems [42], it then follows that $S_0(t) \to S_0^0$ and $S_1(t) \to S_1^0$. So $\mathcal{E}^0$ is globally attractive when $\mathcal{R}_s < 1$. It follows that $\mathcal{E}^0$ is globally asymptotically stable when $\mathcal{R}_s < 1$.

Define:

\[X_0 = \{(S_0, S_1, A, I_s, I_c) \in \mathbb{R}_+^5 : A > 0, I_s > 0, I_c > 0\}, \quad \partial X_0 = \mathbb{R}_+^5 \setminus X_0.\]

Let $P : \mathbb{R}_+^5 \to \mathbb{R}_+^5$ be the Poincaré map associated with system (10) such that $P(x_0) = u(\omega, x_0), \quad \forall x_0 \in \mathbb{R}_+^5$,

where $u(t, x_0)$ denotes the unique solution of the system (10) with $u(0, x_0) = x_0$. It is easy to verify that

\[P^m(x_0) = u(m\omega, x_0), \quad \forall m > 0,\]

Lemma 2.4. When $\mathcal{R}_s > 1$, then there exists a $\delta > 0$ such that when

\[||\psi_x^0, \psi^0, S_0^0, I_0^0 - P_0|| \leq \delta\]

for any $(S_0^0, I_0^0, A_0, S_1^0, I_c^0) \in X_0$, we have

\[
\limsup_{m \to \infty} d[P^m(S_0^0, I_0^0, A_0, S_1^0, I_c^0), P_0] \geq \delta
\]

(16)

where $P_0 = (S_0^0, 0, 0, S_1^0, 0)$. 

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Proof. If \( R_0 > 1 \), we obtain \( \rho(\Phi_{(F - M)}(\omega)) > 1 \) by Lemma 2.3. Choose \( \bar{\epsilon} \) small enough such that
\[
M_{\bar{\epsilon}} = \begin{bmatrix}
\bar{\epsilon} & \bar{\epsilon} & \bar{\epsilon} \\
\bar{\epsilon} & \bar{\epsilon} & \bar{\epsilon} \\
0 & 0 & 0
\end{bmatrix}.
\]

Now we proceed by contradiction to prove that
\[
\limsup_{m \to \infty} d[P^m(S^0, I^0, A^0, S^0, I^0), P_0] \geq \delta.
\]

If not, then
\[
\limsup_{m \to \infty} d[P^m(S^0, I^0, A^0, S^0, I^0), P_0] < \delta
\]

for some \((S^0, I^0, A^0, S^0, I^0) \in X_\delta\). Without loss of generality, we assume that \( d[P^m(S^0, I^0, A^0, S^0, I^0), P_0] < \delta \) for all \( m \geq 0 \). By the continuity of the solution with respect to the initial values, we obtain
\[
\|u(t, P^m(S^0, I^0, A^0, S^0, I^0)) - u(t_1, P_{0})\| \leq \bar{\epsilon}, \quad \forall m \geq 0, \quad \forall t_1 \in [0, \omega].
\]

For any \( t \geq 0 \), let \( t = m\omega + t_1 \), where \( t_1 \in [0, \omega] \) and \( m = \lfloor \frac{t}{\omega} \rfloor \), which is the greatest integer less than or equal to \( \frac{t}{\omega} \). Then we have
\[
\|u(t, (S^0, I^0, A^0, S^0, I^0)) - u(t_1, P_{0})\| \leq \bar{\epsilon}
\]

for any \( t \geq 0 \), which implies that \( S^0 - \bar{\epsilon} < S_0(t) < S^0 + \bar{\epsilon}, S^0 - \bar{\epsilon} < S_0(t) < S^0 + \bar{\epsilon}, \quad t \geq 0 \). Then for \( \|((S^0, I^0, A^0, S^0, I^0) - P_0)\| \leq \delta \), we have
\[
\begin{align*}
\dot{I}_b(t) & \geq \left| \beta_{\omega,\omega}(t)(I_b + \epsilon A) + \beta_{\omega,\omega}(t)I_e[S^0(t) - \bar{\epsilon}] - [\mu_b + \gamma]I_b, \\
\dot{\bar{A}}(t) & = f\gamma I_b - [\mu_b + d_b]A(t), \\
\dot{L}_e(t) & \geq \left| \beta_{\omega,\omega}(t)(I_b + \epsilon A) + \beta_{\omega,\omega}(t)I_e[S^0(t) - \bar{\epsilon}] - [\mu_e + d_e]I_e(t).
\end{align*}
\]

Next we consider the linear system
\[
\begin{align*}
\dot{\bar{I}}_b(t) & = \left| \beta_{\omega,\omega}(t)(I_b + \epsilon A) + \beta_{\omega,\omega}(t)I_e[S^0(t) - \bar{\epsilon}] - [\mu_b + \gamma]I_b, \\
\dot{\bar{A}}(t) & = f\gamma I_b - [\mu_b + d_b]A(t), \\
\dot{\bar{L}}_e(t) & = \beta_{\omega,\omega}(t)(I_b + \epsilon A) + \beta_{\omega,\omega}(t)I_e[S^0(t) - \bar{\epsilon}] - [\mu_e + d_e]I_e(t).
\end{align*}
\]

Once again by Lemma 2.3, it follows that there exists a positive \( \omega \)-periodic function \( \tilde{g}(t) \) such that \( g(t) = e^{pt}\tilde{g}(t) \) is a solution of system (18), where \( p = \frac{1}{2} \ln \rho(\Phi_{(F - M)}(\omega)) \). Because \( \rho(\Phi_{(F - M)}(\omega)) > 1 \), when \( g(0) > 0 \), \( g(t) \to \infty \) as \( t \to -\infty \). Applying the comparison principle [41], we know that when \( I_b(t) \to 0, A(t) \to 0 \) and \( I_e(t) \to 0, I_b(t) \to \infty, A(t) \to \infty \) and \( I_e(t) \to \infty \) as \( t \to -\infty \). This is a contradiction. This completes the proof.

\[\square\]

**Theorem 2.5.** If the basic reproduction number \( R_0 < 1 \), then the unique DFE is globally asymptotically stable in \( \Gamma \). Further, if \( R_0 > 1 \) the disease persists.

### 2.3 Optimal control

In this section, we turn to an optimal control study of our brucellosis models, with an aim of exploring effective prevention and intervention strategies that could best balance the outcomes and costs of the control. To that end, we will perform the optimal control study both the autonomous model (1) and non-autonomous model (10). We introduce two time-dependent control strategies, \( u_1(t) \) and \( u_2(t) \) which are represented as functions of time and assigned reasonable upper and lower bounds. The control function \( u_1(t) \) measures the rate at which susceptible cattle are vaccinated.
during each time period, while control function \( u_2(t) \) accounts for the impact of detection and culling of infectious cattle. Since humans do not transmit the disease we did not consider time dependent intervention strategy. Retaining the same variable and parameter names as in (1), the system of differential equations describing our model with controls is:

\[
\begin{align*}
\frac{dS}{dt} &= \Lambda - [\beta_{sh}(I_c + \epsilon A) + \beta_{sh} I_c]S - \mu S, \\
\frac{dI}{dt} &= [\beta_{sh}(I_c + \epsilon A) + \beta_{sh} I_c]S - [\mu + \gamma]I, \\
\frac{dA}{dt} &= f \gamma I - [\mu + \delta]A, \\
\frac{dC}{dt} &= \Lambda_c - [\beta_{ch}(I_c + \epsilon A) + \beta_{ch} I_c]C - [\sigma u_1(t) + \mu_c]C, \\
\frac{dU}{dt} &= [\beta_{ch}(I_c + \epsilon A) + \beta_{ch} I_c]C - [\mu_c + \alpha]U(t) + \delta I_c, \\
\frac{dH}{dt} &= \Lambda_h - \beta_{sh}(I_c + \epsilon A)S_h + \beta_{sh} I_c S_h - \mu_h S_h + \theta I_h, \\
\frac{d\epsilon}{dt} &= \beta_{sh}(I_c + \epsilon A)S_h + \beta_{sh} I_c S_h - (\theta + \mu_h) I_h.
\end{align*}
\]

According to the extended model above, an optimal control problem with the objective function is formulated by

\[
\text{Minimize } J(u_1(t), u_2(t)) = \int_0^T \left[ B I_c(t) + \frac{W_1}{2} u_1^2(t) + \frac{W_2}{2} u_2^2(t) \right] dt. \tag{20}
\]

The objective is to minimize infected cattle population over a finite time interval \([0, T]\) at minimal costs. In equation (20), \( B I_c \) represent weight constant of the infected cattle. In addition, \( W_1 \) and \( W_2 \) are weight for cattle vaccination and cattle culling. The control efforts in equation (20) are assumed to be nonlinear-quadratic, since a quadratic structure in the control has mathematical advantages, such as: if the control set is compact and convex it follows that the Hamiltonian attains its minimum over the control set at a unique point [43, 44, 45, 46, 47, 48]. Further, \( W_1 u_1^2(t) \), and \( W_2 u_2^2(t) \) describe the costs associated with vaccination and culling, respectively. We assumed that the costs are proportional to the square of the corresponding control function.

The control set is defined as

\[
\Omega = \left\{ (u_1(t), u_2(t)) \mid 1 \leq u_1(t) \leq U_1, 1 \leq u_2(t) \leq U_2 \right\}, \tag{21}
\]

where \( U_1 \) and \( U_2 \) denote the upper bounds for the efforts of vaccination, culling and human treatment, respectively. The bounds reflect practical limitation on the maximum rate of control that can be implemented in a given time period. If, however, \( u_1(t) = u_2(t) = 1 \) for all \( t \), then the model (19) is reduced to the original model (1) or (10), with regular (i.e., minimum) controls.

The optimal control problem hence becomes that we seek optimal functions, \((u_1^*(t), u_2^*(t))\), such that

\[
J(u_1^*(t), u_2^*(t)) = \min_{\Omega} J(u_1(t), u_2(t)) \tag{22}
\]

subject to the state equations in system (19) with initial conditions. The existence of optimal control follows from standard results in optimal control theory [48, 49]. The necessary conditions that optimal controls must satisfy are derived using Pontryagin’s Maximum Principle [50]. Thus, system (2) is converted into an equivalent problem, namely the problem of minimizing the Hamiltonian \( H \)
given by:

\[
H(t) = BL_c(t) + \frac{W_1}{2} u_1^2(t) + \frac{W_2}{2} u_2^2(t) + \lambda_{S_1}(t) \left[ \lambda_c - (\beta_{bc}(I_b + \epsilon A) + \beta_{bh}I_c)S_b - \mu_b S_b \right] \\
+ \lambda_{I_1}(t) \left[ (\beta_{bc}(I_b + \epsilon A) + \beta_{bh}I_c)S_b - (\mu_b + \gamma)I_b \right] \\
+ \lambda_A(t) \left[ f\gamma I_b - (\mu_b + db)A \right] \\
+ \lambda_{S_2}(t) \left[ \lambda_c - (\beta_{bc}(I_b + \epsilon A) + \beta_{cc}I_c)S_c - (\sigma u_1(t) + \mu_c)S_c \right] \\
+ \lambda_{I_2}(t) \left[ (\beta_{bc}u_2(t)(I_b + \epsilon A) + \beta_{cc}I_c)S_c - (\mu_c + \alpha u_2(t) + \delta)I_c \right] \\
+ \lambda_{S_3}(t) \left[ \lambda_c - \beta_{bc}(I_b + \epsilon A)S_b - \beta_{cc}I_cS_b - (\epsilon A + \mu_b + \beta_{bc}I_c) \right] \\
+ \lambda_{I_3}(t) \left[ \beta_{bc}(I_b + \epsilon A)S_b + \beta_{cc}I_cS_b - (\theta + \mu_b)I_b \right],
\]

where \( \lambda_{S_1}(t), \lambda_{I_1}(t), \lambda_A(t), \lambda_{S_2}(t), \lambda_{I_2}(t), \lambda_{S_3}(t) \) and \( \lambda_{I_3}(t) \) denote the adjoint functions associated with the states \( S_b, I_b, A, S_c, \) and \( I_c, \) respectively. Note that, in \( H(t) \), each adjoint function multiplies the right-hand side of the differential equation of its corresponding state function. The first term in \( H(t) \) comes from the integrand of the objective functional.

Given an optimal control treble \( (u_1^*, u_2^*, u_3^*) \) and corresponding states \( (S_b, I_b, A, S_c, I_c) \), there exist adjoint functions \([?]\) satisfying

\[
\frac{d\lambda_{S_1}(t)}{dt} = \frac{dH}{dS_b} = \frac{dH}{dI_b} = \frac{dH}{dA} = -\frac{\partial H}{\partial I_c} = \frac{dH}{dS_c} = \frac{dH}{dI_c} = -\frac{\partial H}{\partial I_b} - \frac{\partial H}{\partial I_c}
\]

These yield

\[
\frac{d\lambda_{S_1}(t)}{dt} = \lambda_{S_1}(t) \left( \beta_{bc}(I_b + \epsilon A) + \beta_{bh}I_c + \mu_b \right) - \lambda_{I_1}(t) \left( \beta_{bc}(I_b + \epsilon A) + \beta_{hh}I_c \right)
\]

\[
\frac{d\lambda_{I_1}(t)}{dt} = \lambda_{S_1}(t) \beta_{bc}S_b + \lambda_{I_1}(t) \left( \mu_b + \gamma - \beta_{bc}S_b \right) - f\gamma A(t) + \lambda_{S_1}(t)\beta_{bc}S_b - \lambda_{I_1}(t)\beta_{bc}S_b + \lambda_{A}(t) \left( \beta_{bc}S_b \right)
\]

\[
\frac{d\lambda_{A}(t)}{dt} = \lambda_{S_1}(t) \beta_{bc}S_b + \lambda_{S_2}(t)\beta_{bc}S_c - \lambda_{I_1}(t)\beta_{bc}S_c - \lambda_{I_1}(t)\beta_{bc}S_b + \lambda_{A}(t) \left( \beta_{bc}S_b \right)
\]

\[
\frac{d\lambda_{S_2}(t)}{dt} = \lambda_{S_1}(t) \beta_{bc}(I_b + \epsilon A) + \beta_{cc}I_c + \mu_c + \sigma u_1(t)(t) - \lambda_{I_2}(t) \left( \beta_{bc}(I_b + \epsilon A) + \beta_{cc}I_c \right)
\]

\[
\frac{d\lambda_{I_2}(t)}{dt} = -B + \lambda_{S_1}(t) \beta_{bc}S_b + \lambda_{S_2}(t)\beta_{bc}S_c + \lambda_{I_1}(t) \left( \mu_c + \alpha u_2(t) + \delta - \beta_{cc}S_c \right)
\]

\[
\frac{d\lambda_{S_3}(t)}{dt} = \lambda_{S_1}(t) \beta_{bc}(I_b + \epsilon A) + \beta_{cc}I_c + \mu_b + \lambda_{A}(t) \left( \beta_{bc}S_b + \beta_{cc}I_c \right)
\]

\[
\frac{d\lambda_{I_3}(t)}{dt} = \lambda_{I_2}(t) \left( \theta + \mu_b \right) - \lambda_{S_3}(t) \theta
\]

with transversality conditions \( \lambda_i(T) = 0 \), for \( i = S_1(t), I_1(t), A(t), S_2(t), I_2(t), S_3(t), I_3(t) \). Furthermore, the optimal controls are characterized by the optimality conditions:

\[
u_1(t) = \max[1, \min(u_1(t), U_1)], \quad u_2(t) = \max[1, \min(u_2(t), U_2)].
\]
where
\[
\begin{align*}
\bar{u}_1(t) &= \frac{\sigma S_{bc} \lambda_{bc}}{2W_1}, \\
\bar{u}_2(t) &= \frac{\alpha L_{bc} \lambda_{bc}}{2W_2}.
\end{align*}
\] (25)

While it is known that wildlife can be important in brucellosis transmission dynamics, lack of data present an enormous challenge to models on evaluating animal and public health control strategies. Wildlife present a complex component of transmission that can be difficult to characterize and there is a need for surveillance data to be coupled with molecular, genetic, and dynamical modeling tools in order to begin to unravel this complexity [7]. Despite the unavailability of surveillance data we proceed to explore the numerical solutions to our autonomous model. Parameter values and variables used in our simulations are listed in Table 2.

In the formulation above, the parameters \( \beta_{ij} \), \( i, j = b, c, h \) can be either constants, for the autonomous model (1), or periodic functions in the form of equation (11), for the periodic model (10). For each case, the state equations, adjoint equations and optimality conditions constitute an optimal control problem, which is then solved numerically.

We adopted the following initial population levels from [13], \( S_c(0) = 1.33 \times 10^6 \), \( L_c(0) = 3.3 \times 10^5 \), \( S_b(0) = 1.618 \times 10^6 \), \( I_b = 0 \) and estimate the African buffalo population levels as follows \( S_b(0) = 4.341 \times 10^7 \), \( I_b(0) = 1.33 \times 10^6 \) and \( A(0) = 0 \). For simplicity we set \( B = 1 \). We further assume that vaccination incurs higher costs than those for culling so that \( W_1 = 10^6 \) and \( W_2 = 10^5 \). In addition, we set the control bound of \( u_1(t) \) and \( u_2(t) \) as follows \( U_1 = 20 \) and \( U_2 = 3 \), respectively.

Baseline values for our model parameter have been adopted from various source abound in literature as indicated in Table 2. Since prior studies suggests that the optimal treatment of uncomplicated brucellosis should be based on a six-week regimen of doxycycline combined either with streptomycin for 2-3 weeks, or rifampicin for six weeks [29], we assume that the average treatment duration is 4 weeks and then \( \theta = \frac{4}{2} = 2 \).

### Table 2. Parameters and values

| Symbol          | Definition                                         | Value                | Units       | Source |
|-----------------|----------------------------------------------------|----------------------|-------------|--------|
| \( (A_b, A_c, A_h) \) | Recruitment rate                                   | (168,000, 197,600, 91,500) | year \(^{-1} \) | [13]   |
| \( (d_b, d_c) \)    | Natural elimination rate                           | (0.04, 0.22, 0.02)   | year \(^{-1} \) | [12, 51] |
| \( a_k \) \( (k = 1, 2, 3, 4) \)   | Amplitude of oscillation                           | 0.8                  |             |        |
| \( \beta_{bc}, \beta_{gh} \) | Averaged direct transmission rate                  | 0.135 \times 10^{-6} | animal \(^{-1} \) year \(^{-1} \) | [13]   |
| \( \beta_{hb} \) | Averaged direct transmission rate                  | 0.21 \times 10^{-6}  | animal \(^{-1} \) year \(^{-1} \) | [13]   |
| \( \beta_{hc} \) | Averaged direct transmission rate                  | 0.18 \times 10^{-6}  | animal \(^{-1} \) year \(^{-1} \) | [13]   |
| \( \beta_{bh} \) | Averaged direct transmission rate                  | 1.3458 \times 10^{-9} | animal \(^{-1} \) year \(^{-1} \) | [13]   |
| \( f \)            | Proportion of new infections that develop into chronic | 0.5896 \times 10^{-9} | animal \(^{-1} \) year \(^{-1} \) | [13]   |
| \( \gamma \)   | Rate of progression to chronic carrier state African buffaloes | 0.67                 | year \(^{-1} \) | [51]   |
| \( \theta \)       | Human treatment rate                               | 13                   | year \(^{-1} \) | [29]   |
| \( \epsilon \)    | Modification factor                                | 0.5                  |             |        |
| \( \sigma \)       | Vaccination rate                                   | 0.316                | year \(^{-1} \) | [11]   |

Fig. 6 depict the numbers of infected wildlife, cattle and human over finite time interval in the presence and absence of optimal control. The results demonstrate that optimal control strategy has a significant effect on the numbers of infected wildlife, cattle and humans, though, these control strategies are targeting cattle population only. Similar behavior is also present on the results for the periodic model (see Fig. 7, but with oscillatory patterns indicate the influence of seasonality on brucellosis dynamics. Results in both Fig. 6 and Fig. 7 shows that the implementation of time dependent intervention strategies aimed targeting domesticated livestock may not be sufficient to
Fig. 6. The numbers of infected wildlife, cattle and human for the autonomous model (1): (a) clinically infected buffaloes (b) chronically infected buffaloes; (c) infected cattle; (d) infected humans.
Fig. 7. The numbers of infected wildlife, cattle and human for the periodic model (10): (a) clinically infected buffaloes (b) chronically infected buffaloes; (c) infected cattle; (d) infected humans.
control the spread of human brucellosis in communities where cattle, wildlife and human interact. However, when there is extremely minimum interaction between wildlife and both cattle and humans (we set $\beta_{cb} = \beta_{bc} = 0.21 \times 10^{-10}$) we observe that optimal vaccination and culling of cattle will lead to brucellosis elimination (see Fig. 8).

Fig. 9 depicts the optimal control profiles for $u_1(t)$ and $u_2(t)$ for the periodic model (10). As we can observe, both $u_1$ and $u_2$ start at the maximum and remain there for a period of 18 years. Thereafter both controls ($u_1$ and $u_2$) begin to oscillate with time for all the remaining period. These results suggest a maximum effort for vaccination and culling for the entire horizon. A similar remark can be drawn for an autonomous model (1) since both controls $u_1$ and $u_2$ start at the maximum and remain there for the entire time horizon see Fig. 10.

To explore the effects of costs on the implementation of control strategies, we varied $W_1$ and $W_2$. Suppose $W_1 = 10^{10}$ and $W_2 = 10^5$. Then as illustrated in Fig. 11 control $u_1$ and $u_2$ will not stay
at the maximum for the entire time horizon as we have observed earlier. More specifically, \( u_1 \) will stay at the maximum for a period of 2 years while \( u_2 \) will stay at the maximum for about 7 years and this is due to the relatively lower value of \( W_2 \).

3 Concluding Remarks

In this paper, we constructed a theoretical framework to investigate the transmission dynamics of human brucellosis in periodic and non-periodic environments. We also investigated the implications of intervention strategies on controlling the spread of brucellosis. To explore brucellosis dynamics in non-periodic environments we developed an autonomous dynamical system with constant parameters that account for all the essential biological dynamics of brucellosis. Our model incorporated human population, cattle population and wildlife population (African buffaloes). We conducted thorough analysis of the model, including computation of the basic reproduction number and stability analysis of the model steady states. Particularly, we demonstrated that when the basic reproduction number
is less than unity then our autonomous model has globally stable disease free equilibrium. This implies that the disease dies out in the community. However, if the basic reproduction number is greater than unity then there exists a unique endemic equilibrium that is globally asymptotically stable.

Extensive investigation of the relationship between reproduction number $R_0$ and model parameters, we observed that model parameters such as $\Lambda_b$, $\Lambda_c$, $\beta_{bc}$, $\beta_{bb}$, $\epsilon$, and $f$, have a positive influence on the $R_0$, that is, whenever they are increased, the size of $R_0$ increases. For example, an increase in recruitment rate of Buffalo $\Lambda_b$ by 73.99% will lead to an increase in the size of $R_0$ by 73.99%. In the other-hand, model parameters with negative index values have a negative influence on $R_0$, for example, an increase in mortality rate of Buffalo $d_b$ by 46.15% will lead to a decrease on the magnitude of $R_0$ by 46.15%. Overall, we noted that increase on progression rate of infected cattle to chronic stage reduce the size of $R_0$. In particular, one can note that whenever the progression rate is greater than 0.5 the disease dies in the community. Furthermore, we observed that whenever the vaccination rate of susceptible cattle is less than 0.5 the disease persists in the community.

The model allowed us to demonstrate the implications of time dependent controls. Specifically we have shown that time dependent intervention strategies for cattle population could minimized brucellosis incidence in both humans and wildlife. In addition, our optimal control simulations demonstrate that if humans and cattle have extremely minimal interaction then brucellosis can be effectively controlled in both humans and cattle and it may require a period of 5 years to successful contain the disease. However, if the intervention strategies are regular the disease may not be successful contained even if the strategies are implemented for a time period of 40 years.

In the second model, we extended the autonomous system to a periodic environment to analyze the impacts of seasonal variation that may affect the movements of animals and, consequently, brucellosis transmission. We derived an expression for the seasonal-induced basic reproductive number and showed that the basic reproductive number remains a sharp threshold for brucellosis dynamics even in a periodic environment. Thus, if the basic reproduction number is less than unity brucellosis will be eradicated. We also proved uniform persistence of the disease as well as the existence of a nontrivial periodic solution when the basic reproduction number is greater than unity. In a similar manner, to our autonomous model, we explored the implication of time dependent cattle vaccination and culling of infected cattle. Our optimal control simulations in this case concurred with our earlier findings on the autonomous model, that optimal control can greatly reduce brucellosis incidence among human, wildlife and cattle. However, our optimal control simulations of the periodic model also exhibited annual oscillations which reflect the effect of seasonality on brucellosis dynamics in periodic environments. In conclusion, our study demonstrate that, in all scenarios, the optimal control can greatly reduce the burden of brucellosis in the community and most importantly if wildlife and cattle and humans and cattle have extremely minimal contacts then the disease can be effectively controlled in a shorter period of time.

A potential limitation of the present paper is that we have employed the mass action incidence for the direct transmission route for both the autonomous and time-periodic model. Such mass action forms, with the advantage of making model calculations more tractable, have been extensively used in previously published brucellosis modeling work (see, e.g., [12, 11, 16, 18]). On the other hand our model did not include indirect transmission, for realistic applications, the indirect (i.e., environment-to-host) transmission it would have been better if it is included in our model and represented by saturated type functional responses, and such type of incidence forms have been used in modeling some other environmentally transmitted diseases (such as cholera [52, 53]). It would be interesting to employ saturated type incidence in our future work on brucellosis modeling, for both the autonomous and time-periodic cases.
Our study can be extended by assessing the impact of intervention strategies that minimize cattle and wildlife interactions such as maintenance of game fencing. Also the study can be strengthened by incorporating heterogeneous interaction between cattle, wildlife and humans. It is also undeniable that fitting those key model parameters with realistic seasonal data will improve our model and its applicability.

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**Competing Interests**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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