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ORIGINAL RESEARCH ARTICLE



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# Budget Impact Analysis of Using Dihydroartemisinin– Piperaquine to Treat Uncomplicated Malaria in Children in Tanzania

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#### Abstract

*Background and Objective* Dihydroartemisinin–piperaquine (DhP) is a very cost effective anti-malarial drug. The aim of this study was to predict the budget impact of using DhP as a first- or second-line drug to treat uncomplicated malaria in children in Tanzania.

*Methods* A dynamic Markov decision model was developed based on clinical and epidemiological data to estimate annual cases of malaria in children aged under 5 years. The model was used to predict the budget impact of introducing DhP as the first- or second-line anti-malarial drug, from the perspective of the National Malaria Control Program in 2014; thus, only the cost of drugs and diagnostics were considered. Probabilistic sensitivity analysis was performed to explore overall uncertainties in input parameters. *Results* The model predicts that the policy that uses artemether–lumefantrine (AL) and DhP as the first- and second-line drugs (AL + DhP), respectively, will save about US64,423 per year, while achieving a 3 % reduction in the number of malaria cases, compared with that of

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AL + quinine. However, the policy that uses DhP as the first-line drug (DhP + AL) will consume an additional \$US780,180 per year, while achieving a further 5 % reduction in the number of malaria cases, compared with that of AL + DhP.

*Conclusion* The use of DhP as the second-line drug to treat uncomplicated malaria in children in Tanzania is slightly cost saving. However, the policy that uses DhP as the first-line drug is somewhat more expensive but with more health benefits.

# **Key Points for Decision Makers**

Understanding the financial burden that will be imposed by a new health technology on the health system is important for planning and budgeting.

Dihydroartemisinin–piperaquine is relatively more expensive than artemether–lumefantrine but has a greater potential to reduce the burden of *Plasmodium falciparum* malaria when used as the first-line drug to treat uncomplicated malaria.

# **1** Introduction

Malaria is an infectious disease, usually of short duration; there were an estimated 198 million cases of malaria (range 124–283 million) and an estimated 584,000 deaths (range 367,000–755,000) globally in 2013 [1]. In Tanzania, malaria is responsible for about one-tenth of all outpatient fevers in children [2]. Most malaria infections in sub-Saharan Africa are caused by *Plasmodium falciparum*, and without adequate treatment, the disease can progress rapidly to life-threatening severe malaria. There is a lot of controversy surrounding the precise burden of malaria in African countries. In 2010, malaria deaths in Tanzania among children aged under 5 years were estimated to range between 10,928 and 49,663 [3]. In 2013, WHO estimates showed that there were about 14.6 million cases of suspected malaria in the country, which places Tanzania among 18 countries with the highest burden of malaria in the sub-Saharan African region [1].

Tanzania has repeatedly changed its malaria treatment guidelines in response to *P. falciparum* resistance to antimalarial drugs. In 2001, chloroquine was replaced with sulfadoxine–pyrimethamine (SP) as the first-line drug to treat uncomplicated malaria. In 2007, SP was replaced with artemether–lumefantrine (AL), while quinine was a recommended second-line drug (AL + quinine) [4]. In 2014, quinine was replaced with dihydroartemisinin–piperaquine (DhP) as the second-line drug (AL + DhP) [5]. AL and DhP are two of the five artemisinin-based combination therapies (ACTs) recommended by the WHO to treat uncomplicated malaria [6].

Recent research evidence indicates that DhP may be a better first-line drug to treat uncomplicated malaria than AL because it is more effective [7] and offers a prolonged post-treatment prophylaxis, which reduces the risk of recurrent malaria infections [8, 9]. In addition, DhP has a simple once-a-day dosage regimen which enhances adherence to treatment [10]. Economic evaluation studies have also indicated that DhP is more cost effective than AL when used as the first-line drug; hence, it represents better value for money [8, 11, 12]. Mori et al. [12] showed that DhP was more cost effective than AL from a providers' perspective, with an incremental cost-effectiveness ratio of \$US12.40 per disability-adjusted life-year (DALY) averted in a Tanzanian setting, based on a valuation performed in December 2012 [12].

Cost-effectiveness analysis compares costs and outcomes of competing interventions, and is an important criterion for priority-setting decisions. However, such analyses fail to report information that can be used to assess whether costeffective interventions are also affordable to the purchaser. Budget impact analysis (BIA), which addresses the issue of affordability, is therefore increasingly used to complement cost-effectiveness analyses. BIA estimates the financial consequences of adopting a new drug within a specific healthcare setting to guide formulary listing decisions [13]. This study aims to predict the budget impact of adopting DhP as the first- or second-line drug to treat uncomplicated malaria in children in Tanzania compared with a previous policy composed of AL and quinine.

# 2 Methods

# 2.1 Analytical Framework

The BIA was conducted using a dynamic Markov cohort model with four mutually exclusive health states: "well", "uncomplicated malaria", "severe malaria", and "death" as an absorbing health state. The model is a modified version of the one used to compare the cost effectiveness of AL and DhP when used as the first-line drugs in Tanzania [12]. While the original model was closed, this is an open model, which allows entrance of new members through births while others exit after reaching the age of more than 5 years over the 1-year time horizon of the model (Fig. 1). The cohort begins in a "well" state and then transits to other health states in 1-week cycles based on risk factors for malaria, access to healthcare, and the effectiveness of anti-malarial drugs.

The analysis was performed from the perspective of the National Malaria Control Program and has a duration of 1 year. A probabilistic sensitivity analysis (PSA) was conducted using Monte-Carlo simulations to assess the robustness of the model considering the uncertainty associated with the input parameters. The analysis was run using TreeAge Pro 2015 software (TreeAge Software, Inc., Williamstown, PA, USA). The study adheres to the BIA guidelines regarding principles of good practice from the International Society for Pharmacoeconomics and Outcome Research (ISPOR) [13].

# 2.2 Intervention Mix

Two competing treatment policies for uncomplicated malaria, each consisting of AL and DhP as first- or secondline drugs, were evaluated against a reference policy of AL + quinine. A regimen consisting of a 3-day dosage of parenteral quinine, followed by an oral dose of the first-line drug, was employed as a standard treatment for severe malaria in all policies. The treatment policies involved are:

- AL + quinine: this is a reference policy, which uses AL and quinine as first- and second-line drugs, respectively. This policy was launched in 2006 and followed until it was replaced in 2014 by AL + DhP [4].
- AL + DhP: this policy option substitutes quinine with DhP as the second-line drug in the reference policy and is consistent with the policy change of 2014 in the country [5]. This treatment policy is also consistent with WHO recommendations regarding the treatment of uncomplicated malaria [6].
- 3. DhP + AL: this policy uses DhP as the first-line drug and AL as the second-line drug. This policy would

Fig. 1 State transition diagram of the model



exploit the benefits of DhP, which include relatively higher efficacy and compliance rates. It is also consistent with WHO recommendations regarding the treatment of uncomplicated malaria [6].

# 2.3 Patient Population

The study was conducted in a sub-population of 7,273,832 children aged under 5 years in Tanzania [14]. Children are most vulnerable to malaria and account for more than twothirds of all malaria deaths [1]. Estimates of the size of the eligible population was based on a 2-week prevalence of fevers of about 20 % [15], of which about 10.5 % have been attributed to malaria infections [2]. The cohort grows at a rate of about 8466 children per week, which is the difference between the birth and aging rate [14, 16]. The all-cause mortality rate of 81 per 1000 live births for children aged under 5 years [17] was adjusted downward by 11 % to account for deaths due to causes other than malaria [18]. Progression to severe malaria and survival rates in each arm depend on the effectiveness of the firstand second-line drugs, which in this study are influenced by differences in compliance rates.

#### 2.4 Characteristics of Anti-Malarial Drugs

A large head-to-head, multicenter randomized clinical trial indicated that DhP has a cure rate of 97.3 versus 95.5 % for

AL, based on an intention-to-treat analysis. The administration of drugs in this trial was directly observed and children stayed at the facilities long enough to check for any vomiting, which implies perfect adherence to treatment [19]. DhP is administered once daily while AL should be taken twice daily together with fat-rich meals, both for 3 consecutive days. Compliance rates for AL and DhP range from 60 to 80 and 70 to 90 %, respectively [10, 20].

Quinine has been used as a second-line drug for the management of uncomplicated malaria in endemic countries for many years, mostly due to the lack of an appropriate alternative drug. The major limitations of quinine are its long, three times daily dosage regimen, which extends to 7 days, and cinchonism adverse effects. In a nested clinical trial, quinine used as a second-line drug to treat recurrent malaria infection in children in Uganda showed cure rates ranging from 88 to 98 % in a head-to-head comparison with AL and DhP [21]. An assumed compliance rate varying from 40 to 60 % was used in this study, based on a clinical trial and one other economic study [22, 23].

Patient compliance to treatment is a primary determinant of therapeutic outcomes. Poor compliance to antimalarial drugs increases the chances of treatment failure and, more importantly, drug resistance. However, in economic modeling studies it is commonly assumed that 10–20 % of non-compliers to multi-dosage anti-malarial drugs such as the ones considered in this study may experience spontaneous recovery [24, 25]. Therefore, in order to predict cure rates, we combined efficacy rates from trial settings and adherence rates observed in routine clinical practice using the following formula (Eq. 1):

$$Cure rate = E_0 C + E_{nc}(1 - C)$$
(1)

where  $E_0$  is efficacy, *C* is the compliance rate, and  $E_{nc}$  is the proportion of non-compliers who may experience spontaneous recovery.

# 2.5 Assumptions

The model was based on the following simplifying assumptions:

- The policy implementation has reached a steady state.
- There is no asymptomatic malaria state; hence, all cases of severe malaria are due to progression from uncomplicated malaria.
- Uncomplicated malaria is not fatal and all positive cases in the public facilities are prescribed the recommended first-line drug.
- The second-line drug ACT will be restricted in the public sector, and when it is out of stock patients will be given the first-line drug.

# 2.6 Care Seeking for Diagnosis and Treatment of Malaria

Care-seeking behavior for the treatment of malaria is very complex and is influenced by many factors, such as perceived severity of the disease, proximity to the facility, availability of medicine, and the ability to pay for health services [26, 27]. In this study, caregivers are modeled to seek care for their febrile children from three main sources: the public sector, which also includes non-profit faithbased facilities; the private sector, which is composed of accredited drug shops and pharmacies; and informal sources. A recent national HIV/AIDS and Malaria Indicator Survey reported that between 77 and 81 % of caregivers sought care from formal health facilities [15]. Between 50 and 70 % of these formal facilities are composed of public facilities, and the remainder include private-sector pharmacies and drug shops, which are scattered all over the country [28].

Between 50 and 70 % of all febrile children who visit public facilities are treated based on malaria Rapid Diagnostic Test (mRDT) results and the rest are treated based on presumptive diagnosis [29, 30]. Non-adherence by health workers to a negative mRDT was estimated to vary from 7 to 14 % [31, 32]. Test results were based on Bayesian calculations, with a prior positive test probability of 10.5 %, and sensitivity and specificity (range) of the mRDT of 95.4 % (94.2–96.6 %) and 95.9 % (94.8–97.0 %), respectively [33]. The sensitivity and specificity (range) of presumptive diagnosis were set at 30 % (20–40 %) and 90 % (80–100 %), respectively [34].

Availability of the first-line drug in public facilities varies from 40 to 80 % [35], and when it is out of stock between 40 and 60 % of patients will be able to access it from private-sector drug shops and pharmacies [36]. The rest may purchase non-recommended drugs, with cure rates ranging between 10 and 60 % [37]. If a child still has uncomplicated malaria after the initial treatment, we assume the caregiver will choose to return to the public facility, where the second-line drug will be prescribed, with availability of 80–100 % for quinine (i.e., because it is relatively cheap) and 40–60 % for ACTs.

Private-sector drug shops and pharmacies are an important first point of care for patients with suspected malaria infections in Tanzania. However, diagnostic tests are usually unavailable; hence, treatments are based on symptoms alone. A majority of these premises stock ACTs, following the implementation of the Affordable Medicines Facility-malaria (AMFm) program [38]. Considering that the major incentive of these premises is to maximize sales, we assume that 60–80 % of febrile children receive antimalarial drugs [39], of which about 40–60 % are a recommended first-line drug [40]. To capture the complexity of care seeking, we assumed that half of the patients who fail to respond to treatments will choose to shift to the public health facilities and the rest will remain in the private sector.

In Tanzania, only formal healthcare facilities, including the private-sector accredited drug shops and pharmacies, are authorized to stock anti-malarial drugs. Therefore, all treatments sought from informal sources such as markets, local shops, and traditional healers are considered to be ineffective. However, we assumed that 10–20 % of these cases may experience spontaneous recovery [12]. Those who continue to suffer from uncomplicated malaria will shift to the public healthcare facilities for further treatment.

Between 3 and 7 % of untreated malaria cases, or those who experience treatment failure, progress to severe malaria [25, 41]. Between 80 and 100 % of severe cases of malaria have been estimated to have access to adequate inpatient care [42], which reduces the case-fatality rate, ranging between 45 and 80 % [43], to about 10.9 % [44].

# 2.7 Resource Use and Costs

A costing study conducted at an urban district-level hospital exists and this has been reported elsewhere [12]. However, due to its lack of representative cost data for personnel and other items, the current BIA study focuses on expenditures incurred on ACTs, mRDTs, and other drugs used to manage the associated co-morbidities. This is more relevant to the National Malaria Control Program, which is responsible for the development of malaria policies and budgeting for ACTs and mRDTs, which are largely funded by donors. For uncomplicated malaria, the drugs included are the recommended first- and second-line anti-malarials. We included SP also to represent the use of non-recommended drugs, especially from the private-sector premises [15, 28]. Antipyretics such as paracetamol were omitted because they are relatively cheap and hence unlikely to have significant budgetary implications. For severe malaria, the cost of a standard regimen consisting of parenteral quinine, diclofenac, diazepam, hematinics, and a first-line ACT was included [5].

The Global Fund's reference prices negotiated with the manufacturers of ACTs for children aged under 5 years in endemic countries were used [45], while for SP a buyer price (range) of \$US0.32 (\$US0.25–0.38) from the *International Drug Price Indicator Guide* was used [46]. Prices of ACTs were inflated by 20 % to account for program costs, i.e., 10 % freight and insurance costs [46] and 10 % as estimated local distribution costs. Prices of other drugs were taken from the price catalogue of the Medical Stores Department (MSD) [47]. Prices of mRDTs have decreased recently due to price competition; hence, a unit price (range) of \$US0.45 (\$US0.36–0.55) was used [30]. An exchange rate of 1670 Tanzanian shillings to \$US1 for the year 2014 was used to convert local MSD prices into US dollars [48].

#### 2.8 Uncertainty

Probability distributions were employed to incorporate uncertainties in the parameters used in the model (Table 1). Beta distributions were used for probability parameters to limit their possible values to the interval 0–1, while costs were constrained between 0 to positive infinity by gamma distributions. Normal distributions were used to describe estimated population size, birth rates, and compliance rates of different anti-malarial drugs. Overall uncertainty in the parameters was propagated in the model by running a PSA based on a Monte-Carlo simulation with 10,000 iterations. One-way sensitivity analysis was used to test how variations in key parameters, including addition of hospitalization costs, may potentially influence the results.

# **3** Results

## 3.1 Model Validation

The model was validated using the reference policy (AL + quinine) to determine whether the predicted health

outcomes from the simulations correspond with the figures reported in the literature. An estimated 10-12 million cases of uncomplicated malaria occur in Tanzania annually, of which about two-thirds are in children aged under 5 years [30]. Our model prediction of 7,510,727 cases is consistent with the implied range therefore of 6.7-8.0 million cases. The model predicted 173,600 cases of severe malaria among children aged under 5 years, which is also about two-thirds of all severe cases based on the WHO World Malaria Report 2013 [50]. The predicted total number of deaths in children aged under 5 years was 134,028, which is within the range of 123,100-186,700 [51]. The predicted number of malaria-attributable deaths was 26,973, which is within the reported range of 10,928–49,663 for children aged under 5 years in Tanzania [3].

#### 3.2 Budget Impact Analysis

The model predicts that the treatment policy of AL + DhPwill save about \$US64,423 (0.3 %) per year, compared with that of AL + quinine, and the policy of DhP + ALwill consume an additional \$US780,180 (3.8 %) per year compared with that of AL + DhP. These represent changes in the budget for drugs and mRDTs only (Table 2). It is interesting to note that DhP + AL has the highest drug costs but the lowest costs for mRDT compared with other policies. The reason for this difference in cost is that the higher effectiveness rate of DhP as a first-line drug reduces episodes of malaria and hence the requirement for mRDT.

# 3.3 Impact on Health Outcomes

Table 3 shows the estimated change in overall annual health outcomes, as recommended by the new guidelines for reporting BIA analysis [13], for the two malaria treatment policies. The model predicts that the policy of AL + DhP has the potential to reduce the number of malaria cases and deaths by 248,437 (3 %) and 1954 (7.2 %), respectively, compared with the reference policy of AL + quinine. However, the policy of DhP + AL, which is more expensive, has the potential to reduce the number of malaria cases and deaths further by 364,517 (5 %) and 2868 (11.5 %), respectively, compared with the policy of AL + DhP.

#### 3.4 Sensitivity Analyses

The policy of AL + quinine has already been replaced with that of AL + DhP, and since it is probably not very attractive to revert to quinine-based therapy, policy makers will be more interested in a comparison of DhP + AL versus AL + DhP. This is presented in the Tornado

Parameters	Estimates	Distributions	Sources
Cohort size of children aged under 5 years	7,273,832 ± 20 %	Normal	[14]
Weekly cohort growth rate	$8466\pm20~\%$	Normal	[14, 16]
Under-five mortality rate per 1000 live births	81 (72–90 %)	Beta	[17]
Malaria-attributed deaths in children aged under 5 years	11 %	Point estimate	[18]
2-Week incidences of fever episodes per child	$0.20$ $\pm$ 20 $\%$	Beta	[15]
Percentage of febrile episodes attributed to malaria	$10.5$ $\pm$ 20 %	Beta	[2]
Case fatality rate of untreated severe malaria	60 (45-80 %)	Beta	[43]
Case fatality rate of treated severe malaria	10.9 %	Beta	[49]
Early treatment failure leads to severe malaria	5 (3-7 %)	Beta	[41]
Untreated malaria becomes severe	5 (3-7 %)	Beta	[25]
Probability of care seeking in formal facilities	79 (77-81 %)	Beta	[15]
Percentage of formal facilities belonging to public sector	60 (50-70 %)	Beta	[28]
Percentage of severe cases with access to inpatient care	90 (80-100 %)	Beta	[42]
Access to first-line drugs in public facilities	60 (40-80 %)	Beta	[35]
Access to second-line drugs in public facilities	50 (40-60 %)	Beta	Assumption
Percentage of children given anti-malarials in private facilities	70 (60-80 %)	Beta	[39]
Percentage of anti-malarials that are ACT in private facilities <sup>a</sup>	50 (40-60 %)	Beta	[36, 40]
Efficacy of DhP	97.3 (94.9–99.7 %)	Beta	[19]
Efficacy of AL	95.5 (93.1-97.9 %)	Beta	[19]
Efficacy of quinine	93.0 (88.0-98.0 %)	Beta	[21]
Compliance with AL	70 (60-80 %)	Normal	[10, 20]
Compliance with DhP	80 (70-90 %)	Normal	[10, 20]
Compliance with quinine	50 (40-60 %)	Normal	[22, 23]
Non-compliers with treatments who recover	20 (10-30 %)	Beta	[24, 25]
Sensitivity of mRDT	95 (94.2-96.6 %)	Beta	[33]
Specificity of mRDT	96 (94.8-97.0 %)	Beta	[33]
Sensitivity of clinical diagnosis	30 (20-40 %)	Beta	[34]
Specificity of clinical diagnosis	90 (80-100 %)	Beta	[34]
Adherence to a negative mRDT result <sup>b</sup>	10.5 (7.0-14.0 %)	Beta	[31, 32]
Drugs and diagnostic costs (\$US per dose/test)			
DhP: Dh 40 mg, P 320 mg (" $3 \times 1$ " pack)	0.77 (0.56-0.93)	Gamma	[45]
AL: A 20 mg, L 120 mg ("6 × 2" pack)	0.67 (0.54-0.84)	Gamma	[45]
Quinine 300 mg/tablet	0.59 (0.47-0.70)	Gamma	[46]
SP: S 500 mg, pyrimethamine 25 mg (" $3 \times 1$ " pack)	0.32 (0.25-0.38)	Gamma	[46]
Quinine injection 300 mg/mL (2 mL ampoule)	2.40 (1.92-2.87)	Gamma	[47]
Diazepam injection 5 mg/mL (2 mL ampoule)	0.23 (0.18-0.27)	Gamma	[47]
Diclofenac injection 25 mg/mL (3 mL ampoule)	0.29 (0.23-0.35)	Gamma	[47]
Dextrose 5 % (500 mL bottle)	3.95 (3.16-4.74)	Gamma	[47]
Ferrous sulphate + folic acid $200 + 0.25$ mg	0.26 (0.21-0.31)	Gamma	[47]
Cost of mRDT	0.45 (0.36-0.55)	Gamma	[30]

Numbers in the parentheses represent ranges

A artemether, ACT artemisinin-based combination therapy, AL artemether-lumefantrine, Dh dihydroartemisinin, DhP dihydroartemisininpiperaquine, L lumefantrine, mRDT Rapid Diagnostic Test for malaria, P piperaquine, S sulfadoxine, SP sulfadoxine-pyrimethamine

<sup>a</sup> As a proportion of dispensed anti-malarial drugs

<sup>b</sup> Proportion of children who will be treated despite having a negative test result

diagram shown in Fig. 2, which expresses the potential influence of uncertain parameters on budget impact. Cost and the compliance rates for DhP and AL were identified to be the most influential parameters.

Considering that the cost of DhP is the most influential parameter, it was important to show how variation in the cost of DhP causes changes in the budget of drugs and diagnostics, as indicated in Fig. 3. At a cost of about **Table 2**Annual incrementalcost from the perspective of theNational Malaria ControlProgram (\$US)

Policy options	mRDT costs	Drug costs	Total cost	Incremental cost
AL + quinine	6,350,032	14,523,023	20,873,055	Reference
AL + DhP	6,325,074	14,483,558	20,808,632	-64,423 (0.3 %)
DhP + AL	6,232,222	15,356,589	21,588,811	780,180 (3.8 %)

AL artemether-lumefantrine, DhP dihydroartemisinin-piperaquine, mRDT Rapid Diagnostic Test for malaria

Table 3 Impact on the number of malaria cases and deaths for children aged under 5 years

Policy options	Uncomplicated malaria	Severe malaria	All cases	Incremental cases (all)	Number of deaths	Incremental deaths
AL + quinine	7,510,727	173,599	7,684,326	Reference	26,973	Reference
AL + DhP	7,274,872	161,016	7,435,888	-248,437 (3.2 %)	25,019	-1954 (7.2 %)
DhP + AL	6,928,818	142,554	7,071,371	-364,517 (5.0%)	22,151	-2868 (11.5 %)

AL artemether-lumefantrine, DhP dihydroartemisinin-piperaquine



\$US0.84 per dose of DhP, the policy of DhP + AL is the cheapest option and requires a total budget of about \$US20.7 million annually. It should be noted that this cost includes 20 % program costs; hence, the actual acquisition cost is \$US0.67 per dose. That is to say, for the purpose of cost containment, the additional budget of about \$US780,180 required by the more effective policy of DhP + AL may be avoided if DhP could be bought at a price lower than \$US0.67 per dose.

Table 4 shows the budget impact of including hospitalization costs of about \$US76.46 (range \$US60–90) per patient from our previous study [12], with the assumption that all severe cases of malaria will be treated in the public facilities. The model predicts that the policy of AL + DhP will save about \$US934,200 (2.9 %) per year compared with that of AL + quinine, and that of DhP + AL will save about \$US500,342 (1.6 %) per year compared with that of AL + DhP. Therefore, the policy that uses DhP as Fig. 3 Change in total budget of drugs and diagnostics versus variation in the cost of dihydroartemisinin– piperaquine. AL artemether– lumefantrine, DhP dihydroartemisinin–piperaquine



Table 4Budget impact afterincluding hospitalization costs(\$US)

Policy options	Drugs and mRDT costs	Hospitalization costs	Total costs	Incremental costs
AL + quinine	20,873,055	11,803,699	32,676,754	Reference
AL + DhP	20,808,632	10,933,923	31,742,555	-934,200 (2.9 %)
DhP + AL	21,588,811	9,653,401	31,242,212	-500,342 (1.6 %)

AL artemether-lumefantrine, DhP dihydroartemisinin-piperaquine, mRDT Rapid Diagnostic Test for malaria

the first-line drug (DhP + AL) is the most cost-saving treatment policy for uncomplicated malaria when hospitalization costs are taken into consideration.

# 4 Discussion

Malaria is an infectious disease, which consumes a substantial portion of the limited health budgets of sub-Saharan African countries. It is the leading cause of morbidity and mortality among outpatient visits and inpatient admissions in Tanzania. It has been estimated that malaria accounts for about 2 % of the gross domestic product (GDP), which is equivalent to about 20 % of the total health expenditure in Tanzania [52]. This is the first study in Tanzania to estimate the budget impact of introducing DhP to treat uncomplicated malaria in children aged under 5 years either as the first- or second-line drug.

A majority of countries in sub-Saharan Africa are using AL as the first-line drug and quinine as second-line drug against uncomplicated malaria. Even though ACTs are the preferred choice for the treatment of uncomplicated *P. falciparum* malaria [6], it is understandable that some countries persist with the use of quinine as the second-line drug due to a lack of alternative ACTs. Therefore, the emergence of DhP as a very promising ACT has changed the treatment dynamics of malaria in many countries in recent years. A number of countries have already adopted DhP as the second-line drug and many others are considering doing the same. Mainland Tanzania is one of the countries that recently replaced quinine with DhP as a second-line drug to treat uncomplicated malaria [5].

Several studies have shown that DhP is more cost effective than AL to treat uncomplicated malaria in African countries [8, 11, 12]. However, the arbitrary thresholds of \$US150 per DALY averted or 1–3 times GDP per capita, which are commonly used to categorize interventions as being cost effective or not, may not necessarily reflect the willingness and ability to pay for a new intervention for any particular country [53, 54]. In reality, a new drug can

be very cost effective but unaffordable and, for this reason, BIA is increasingly being used to complement cost-effectiveness analyses.

This study found that by considering the costs of drugs and diagnostics only, the policy that uses DhP as the second-line drug (AL + DhP) to treat uncomplicated malaria in children is slightly cost saving, i.e., \$US64,423 per year, while reducing cases of malaria by about 3 % compared with the policy of AL + quinine. The model also predicts that the policy that uses DhP as the first-line drug (DhP + AL) increases the budget by about \$US780,180 per year, while reducing the number of malaria cases by a further 5 %, compared with AL + DhP. Pfeil et al. [11] found that the use of DhP as the first-line drug in moderateto-high transmission areas will avert 12 % of malaria cases in children. More recently, Okell et al. [8] estimated a reduction of 10-15 % in high-transmission areas. Our study shows a smaller reduction in malaria cases because we did not consider the longer prophylactic effect of DhP, which reduces the recurrence of malaria [8, 9].

Hospitalization costs are an important component of inpatient care costs incurred by healthcare providers to treat severe childhood malaria. The inclusion of hospitalization costs in the sensitivity analysis switched the most cost-saving treatment policy for uncomplicated malaria to DhP + AL, which eventually saves about \$US500,342 per year compared with AL + DhP. Considering that the policy of DhP + AL reduces the number of malaria cases by 5 versus 3 % with AL + DhP, the inclusion of other outpatient and inpatient costs will make DhP + AL even more cost saving compared with AL + DhP.

This study has a number of limitations and, hence, its results must be interpreted with care; more importantly, it should never be used as the sole basis for initiating policy change in Tanzania. Firstly, policy change is a very complex and expensive undertaking [55]. In 2000, it cost the Tanzanian Government about \$US0.8 million to implement the new malaria treatment policy, representing about 4 % of total malaria expenditure and 1 % of total public expenditure on health [56]. Another study estimated the costs of this policy change to be equivalent to \$US0.2 per person [55].

Secondly, care-seeking behavior for diagnosis and treatment of malaria is very complex in sub-Saharan Africa and data may not always be readily available. In the model, a number of structural and parameter assumptions were made in an attempt to replicate how patients move from one type of facility to another in actual practice when seeking care. It is nearly impossible to model this with sufficient accuracy. The model assumptions portray caregivers seeking care from the informal and formal health facilities in a very orderly manner, which rarely happens in reality. Even the diagnosis and treatment practices in public facilities vary from one location to another.

Thirdly, the study assumes that the majority of patients visiting public facilities are diagnosed with mRDTs and all positive cases are prescribed with the recommended first-line drug, with a mean availability of 60 %. While we believe that these are reasonable base-case assumptions, several studies have shown that quality of care in public facilities varies greatly and that mRDTs and drugs often completely run out of stock for prolonged periods of time [57, 58]. The availability of a second-line drug was assumed to be restricted in the public facilities, which may not be the case still. Dispensing practice in private drug shops and pharmacies is more complex than the way it is modeled here; hence, the estimated health benefits of the two malaria policies may deviate from the model predictions.

Fourthly, this study ignored the prolonged post-treatment prophylactic effect of DhP, which has more potential to prevent recurrent malaria infections than AL [8, 9]. The model shows that an increase in the effectiveness of DhP reduces the number of malaria cases and, hence, treatment costs to the greatest extent when it is used as the first-line drug. The inclusion of a prolonged prophylactic effect would have similar implications, i.e., an increase in health benefits and reduction in treatment costs, which would eventually make the policy of DhP + AL even more attractive. Therefore, this study has underestimated the actual benefits of the treatment policy that uses DhP as the first-line drug.

Finally, but equally importantly, rational social planners are concerned not only about reducing healthcare costs but also about broader considerations when distributing scarce healthcare resources, including considerations of efficient and equitable healthcare.

# 5 Conclusion

In accordance with the present model's predictions, the use of DhP as the second-line drug (AL + DhP) to treat uncomplicated malaria in children in Tanzania is slightly cost saving. However, the policy that uses DhP as the firstline drug (DhP + AL) is somewhat more expensive as it consumes an extra US780,180 per year, which represents a 3.8 % increase in the budget for drugs and diagnostics, but with more health benefits than AL + DhP. Nevertheless, the use of relatively more expensive drugs such as DhP as first-line anti-malaria drugs without proper diagnosis should be approached with caution. Otherwise, the opportunity cost of presumptive treatment outweighs the benefits due to over-diagnosis and over-treatment of patients without clinical malaria. Acknowledgments The contribution by BR and OFN was supported by the Centre for Intervention Science in Maternal and Child Health (CISMAC; Project Number 223269), which is funded by the Research Council of Norway through its Centres of Excellence Scheme and the University of Bergen (UiB), Norway. We also thank the Priority-Setting and Economics Research Group, and the two anonymous reviewers for their constructive comments on early drafts of the manuscript.

Author contributions ATM, OFN, and BR conceived the study. ATM retrieved the data and performed the analysis. ATM and BR prepared and designed the economic model. ATM prepared the first draft of the manuscript. BR and OFN supervised data analysis and contributed to the manuscript writing. All authors read and approved the final manuscript.

#### **Compliance with Ethical Standards**

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