Artemisinin, the Magic Drug Discovered from Traditional Chinese Medicine

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A B S T R A C T
Artemisinin and its derivatives represent the most important and influential class of drugs in the fight against malaria. Since the discovery of artemisinin in the early 1970s, the global community has made great strides in characterizing and understanding this remarkable phytochemical and its unique chemical and pharmacological properties. Today, even as artemisinin continues to serve as the foundation for antimalarial therapy, numerous challenges have surfaced in the continued application and development of this family of drugs. These challenges include the emergence of delayed treatment responses to artemisinins in malaria and efforts to apply artemisinins for non-malarial indications. Here, we provide an overview of the story of artemisinin in terms of its past, present, and future. In particular, we comment on the current understanding of the mechanism of action (MOA) of artemisinins, and emphasize the importance of relating mechanistic studies to therapeutic outcomes, both in malarial and non-malarial contexts.

1. Introduction
Malaria has been a debilitating disease with global influence since ancient times and continues to be one of the most widespread and damaging infectious diseases today [1]. With the cause of disease having long been misattributed to “bad air,” the transmissible and parasitic nature of malaria remained unknown until the works of Charles Louis Alphonse Laveran and Ronald Ross in the late 1800s. Their findings established that protozoa belonging to the genus Plasmodium caused malaria, and that Anopheles mosquitoes were the primary vectors of malarial infections. These observations made Laveran and Ross two of the earliest recipients of the Nobel Prize in Physiology or Medicine [2].

In the decades following their discoveries, ground-breaking progress has been made in the battle against the disease. The crusade launched by the Chinese government in the late 1960s to search for cures for malaria ultimately culminated in the discovery of artemisinin. Artemisinin (and its various derivatives, which we will refer to collectively as “artemisinin” unless otherwise specified) is a sesquiterpene lactone compound (Fig. 1) with a unique chemical structure derived from the sweet wormwood plant, Artemisia annua L. (Fig. 2). Since its discovery, it has become the most important and effective antimalarial drug [3].

In many ways, artemisinin is a truly fascinating drug. From the tumultuous process of its discovery, which was deeply tied to traditional Chinese medicine (TCM), to its remarkable potency and impact as an antimalarial drug, it is not surprising that artemisinin has captured a great deal of attention since its introduction to the world stage [1]. Over 40 years after its discovery, artemisinin remains our bulwark against malaria and is the foundation of all major antimalarial therapies [4]. Years of research spanning a range of disciplines have gone into the exploration and elucidation of the mechanisms of artemisinin in its antimalarial role [5]. Beyond that, efforts have been made to repurpose artemisinin for non-malarial applications, thereby raising considerable anticipation over the future development of this drug [6].

With that in mind, we feel that it is a good time to broadly review the timeline of this influential drug, spanning its past, present, and future. Beginning with a look back at the story of the discovery and development of artemisinin, we then review...
and discuss the contemporary understanding of the mechanism of action (MOA) of artemisinin in malaria. We conclude by looking ahead at current efforts to repurpose artemisinin for possible roles outside of malaria. We believe that this article will provide a well-rounded background of artemisinin, along with relevant insights into the salient topics surrounding this remarkable drug.

2. The journey of discovery

We begin with a brief tracing of the remarkable journey that led to the discovery and development of artemisinin. Records of malaria in TCM date back thousands of years, and the same is true for the usage of *Artemisia* (Qinghao) plants as medicinal herbs. First mentioned as a specific remedy for malarial symptoms in Ge Hong’s *Zhouhou Beiji Fang* (*Handbook of Prescriptions for Emergency*) dating back to the Eastern Jin Dynasty (317–420 AD), the application of Qinghao and other techniques for malarial relief was subsequently noted in a series of historical Chinese medical writings that included the influential *Bencao Gangmu* (*Compendium of Materia Medica*) by Li Shizhen (Ming Dynasty, 1368–1644 AD).

This wealth of ancient knowledge would later prove to be instrumental in the discovery and development of artemisinin.

In the years following World War II, the development and deployment of the potent insecticide dichloro-diphenyl-trichloro-ethane (DDT) and new antimalarial drugs such as chloroquine (CQ) resulted in great progress in combating malaria. However, the World Health Organization (WHO)’s campaign in the 1950s to combat and eradicate malaria around the world was eventually met with challenges related to resistance. The emergence of DDT-resistant vectors and drug-resistant parasites led to a rebound of the disease, especially in regions such as Southeast Asia and sub-Saharan Africa [7]. This setback prompted an urgent need for novel antimalarial drugs. Significant efforts had been made by the United States due to the Vietnam War and the prevalence of drug-resistant malaria in that region. The Chinese government also initiated efforts in malarial research around this time. In particular, a national project called Project 523 (named after its date of inauguration, 23 May 1967) was set up to consolidate malarial research on a national level [8].

In 1969, Professor Youyou Tu was selected to lead a research group within the project that focused on screening TCM for novel antimalarial drugs. This work took place at the Institute of Chinese Materia Medica of the China Academy of Chinese Medical Sciences. Drawing from a massive repository of TCM knowledge that included ancient literature, folklore, and oral interviews with practitioners, Tu and colleagues worked from a list of over 2000 herbal remedies, of which some 640 were deemed to be possible “hits.” From this selection, over 380 extracts from approximately 200 herbs (including Qinghao/Artemisia extracts) were eventually collected and tested, mostly giving unsatisfactory results [1,9]. The Qinghao extract nevertheless drew particular interest starting around 1971, as it produced promising but inconsistent results [1]. This finding prompted a revisitation of the literature, and led to perhaps the most important breakthrough in the discovery process.

Returning to the earliest record of the use of Qinghao to treat malarial symptoms, which was in Ge Hong’s *Zhouhou Beiji Fang*, Tu noted that the instructions for the Qinghao prescription involved consuming the strained “juice” of the Qinghao plant immersed in water. It was notable that the instruction made no mention of heating the medicine—something that was otherwise common for prescriptions in TCM. Drawing from the literature and her own knowledge of TCM, Tu arrived at the idea to modify the extraction process to use low-temperature conditions. The extracts produced from this new procedure were further purified by separation of the acidic and neutral phases in order to retain active components while reducing the toxicity of the original extract. The resultant substance displayed a striking 100% effectiveness against rodent malaria in experiments carried out around October 1971. This remarkable result was then fully reproduced in monkey malaria experiments carried out in late December of the same year, thus establishing the efficacy of the Qinghao extract beyond doubt [1].
The breakthrough had been made, but the journey of drug development was by no means complete. Conditions in China at that time made it difficult to perform clinical trials of new drug candidates to ascertain their safety for humans. In an attempt to accelerate the process due to the seasonal and time-sensitive nature of malarial research, Tu and colleagues decided to volunteer themselves as the first human subjects for toxicity and dose-finding tests [8]. This act established the safety profile of the Qinghao extract and enabled clinical trials to be carried out immediately, in the latter half of 1972. The trials (which were carried out in Hainan Province and at the 302 Hospital PLA (now incorporated into the Fifth Medical Center of the Chinese PLA General Hospital) in Beijing) proved successful, and paved the way for Qinghao research to be pushed to the national level. A subsequent concerted effort on the part of the Chinese scientific community at large drove further research and development of Qinghao forward. The active component of the Qinghao extract, artemisinin (also known as Qinghaosu itself), was isolated in November 1972 by Tu's team at the Institute of Chinese Materia Medica. The team would later go on to develop dihydroartemisinin (DHA), which remains one of the most pharmacologically relevant derivatives today. In collaboration with other institutes across China, further groundwork in drug development, including the determination of the stereo-structure of artemisinin and further derivatization of artemisinin, was carried out in the following decade [10,11]. These efforts, among others, culminated in the fourth meeting of the Scientific Working Group on the Chemotherapy of Malaria held in Beijing in 1981, where the findings were presented by Tu for the first time. The results were published in 1982 as a series of papers under the name “China Cooperative Research Group on Qinghaosu and Its Derivatives as Antimalariais” [12,13]; thus the gift from Chinese medicine was delivered to the rest of the world.

In the subsequent years of the 1980s, artemisinin and its derivatives were successfully employed in China to treat thousands of malaria patients [1]. As the problem of drug-resistant malaria continued to worsen elsewhere, it was not long before the commencement of clinical studies with artemisinin in other endemic regions in Asia [14–19]. Consistent and encouraging results led to the expansion of such studies, particularly toward Africa [19–24]. The evidence was clear that artemisinin-based therapy, especially in combination with a slower-acting antimalarial such as mefloquine or piperaquine, led to significant improvements in parasite clearance and a rapid diminishing of symptoms for both uncomplicated and severe Plasmodium falciparum malaria infection. At the same time, its tolerability was shown to be excellent, as reports of toxicity and safety concerns remained minimal [25]. Across more than a decade’s worth of independent randomized clinical studies and meta-analyses, the outstanding efficacy and safety of artemisinin-based therapy became increasingly clear. Finally, in 2006, the WHO announced an alteration of its strategy to fully employ artemisinin combination therapies (ACTs) as the first-line treatment against malaria [26]. ACTs remain the most effective and recommended antimalarial therapies today [4].

3. The search for a mechanism of action

It has been more than a decade since the implementation of ACTs as the official first-line treatment for malaria and over three decades since the discovery of artemisinin. In this time, the clinical and pharmacological characteristics of artemisinin therapy have been extensively scrutinized and reported [27–30]. Although the specifics of various derivatives can differ, artemisinin drugs are characterized by rapid action and potency, low toxicity, and a short half-life, which makes combination therapy with longer-acting antimalarial drugs ideal and recommended [30]. Apart from its pharmacological properties, elucidating the MOA of a drug is important for optimizing treatment regimens. Dosages, drug combinations, and even considerations of drug resistance are closely related to the molecular basis of a drug’s activity. It is thus somewhat surprising that despite decades of widespread application, our understanding of the MOA of artemisinin remains fairly incomplete. Here, we provide a brief overview of the prevailing understanding as well as more recent developments in mechanistic studies of artemisinin [31,32]. In general, the outstanding therapeutic properties of artemisinin can be thought of as a result of two major processes: its unique mechanism of activation, and its downstream activity and drug targets. These mechanisms combine to yield a highly potent, yet highly specific, drug.

3.1. Drug activation

Artemisinin and its derivatives are sesquiterpene lactones that bear the 1,2,4-trioxane moiety as the pharmacophore [33]. In particular, the endoperoxide bridge within this group is well understood to be essential for the pharmacological activity of artemisinin [13,34,35]. Artremisinins are prodrugs in two senses: first, many derivatives are rapidly converted to DHA in vivo, and second, their MOA depends on activation by cleavage of the endoperoxide bridge. The mechanism of this cleavage remains an issue in active research [36]. Malarial parasites are characterized by extensive hemoglobin uptake and digestion during the erythrocytic stage of their life cycle [37,38]. This releases copious amounts of free redox-active heme and free ferrous iron (Fe2+), which are thought to underlie the parasite specificity of artemisinin. Indeed, hemoglobin digestion has been strongly linked with artemisinin susceptibility in parasites [38,39]. Multiple models have been proposed with regard to the mechanism of endoperoxide cleavage by either free redox-active heme or free ferrous iron, and the downstream molecular events that follow cleavage [36,40–48]. These proposals differ in terms of the nature of the cleavage and the identity of the reactive intermediates produced by drug activation. In general terms, however, they explain the parasite-specific drug activation through which reactive species are produced, leading to cellular damage and parasite killing. Recent evidence suggests that free redox-active heme may play a predominant role in drug activation [49,50]. A 2008 study provided in vitro data that indicated that ferrous heme may be a stronger activator of artemisinin than other iron-containing species, including hemin, free ferrous iron, and undigested hemoglobin [49]. Similar observations were made in live parasites, in which artemisinin activation was blocked by inhibiting hemoglobin digestion but not by the chelation of free ferrous iron [47]. Thus, the process of hemoglobin digestion in infected erythrocytes, which is required for parasite growth, is the key to the specificity of artemisinin activation [38].

Interestingly, in studies using yeast cells as a proxy for malaria parasites [51,52], it was found that mitochondria were directly involved in both the activation and action of artemisinin, thus further linking artemisinin action to reactive oxidative species (ROS) production and oxidative damage. It is also plausible that multiple redundant activation pathways may exist in different environments or localities, where the conditions and magnitude of activation can differ [53]. Looking ahead, it will be crucial to consider the pivotal role of drug activation in the activity of artemisinin and to further elucidate its mechanisms under different conditions.

3.2. Downstream mechanism

The crucial step in elucidating a drug’s MOA is to identify its cellular targets. In the conventional understanding of drug design and mechanisms, a drug modifies one or more specific cellular targets, such as proteins, in order to effect downstream changes. However,
the exceedingly fast-acting and potent nature of artemisinin activity, taken together with its ability to alkylate targets, may be due to quite a different mechanism.

First of all, heme releases from hemoglobin digestion functions that lie beyond drug activation, as previously outlined. Excess heme is converted in infected erythrocytes to hematin, which is toxic to the parasite via oxidative damage and direct lysis of cell membranes [54]. Malarial parasites have therefore evolved a detoxifying mechanism that converts hematin to the nontoxic and inert crystalized hemozoin via a biocrystallization process [55]. Activated artemisinin has been reported to prevent the formation of hemozoin by alkylating heme; therefore, it functions in a similar capacity to other antimalarial drugs that act on hemozoin formation, such as CQ [45,56–58]. Thus, free heme from hemoglobin digestion serves as both the activator and the target of artemisinin [45].

Given that activated artemisinin is thought to generate ROS, it is unsurprising that artemisinin has also been reported to directly alkylate protein targets [59,60]. The translationally controlled tumor protein (TCTP) and the alkylate protein targets [59,60]. The translationally controlled tumor protein (TCTP) and the Plasmodium sarcoendoplasmic reticulum Ca²⁺-ATPase PfATP6 were among the first targets of interest that were identified as interacting partners of artemisinin [61–63]. Consideration of the role of single targets in the activity of artemisinin has now evolved into MOAs that may depend on multiple targets, as later studies have shown [64–67]. Using unbiased proteomics methods, it has been observed that artemisinin targeting may be promiscuous rather than monotarget-specific. In the first study that systematically reported artemisinin binding targets, over 100 proteins were identified in live parasite strains [47]. An independent study carried out by Ismail et al. [68] led to consistent findings. These results support a promiscuous mechanism of artemisinin targeting in which activated artemisinin alkylates and damages many cellular proteins, thereby disrupting multiple key biological functions and resulting in toxicity and lethality in parasites [47,48,50]. Interestingly, PfATP6 and other key transporters such as PfCRT and Pfmdr1 are consistently labeled in these types of experiments. These findings are consistent with PfATP6 being an important target for artemisinins [47,68]. As an independent line of evidence, the mapped binding sites of artemisinin to TCTP further support a heme-activated promiscuous mechanism in which modification sites are proximity-based and essentially random [50].

Our current knowledge of artemisinin paints a picture of a drug with a unique and elegant mechanism. Artemisinin and its derivatives are prodrugs that absolutely require endoperoxide group cleavage for drug activation and subsequent anti-parasitic activity. Artemisinin activation is dependent on a heme-rich environment, which is specific to infected erythrocytes as well as being an unavoidable outcome of parasite metabolism. The heme-rich environment itself is then exploited by the activated drug to achieve efficient parasite killing. This mechanism essentially links infection and parasite growth to drug activation, thus ensuring both the outstanding specificity and the tolerability of artemisinin therapy. At the same time, activated artemisinin indiscriminately damages proximal proteins and cellular structures. Rather than targeting a single protein or cellular function, like the majority of conventional drugs (including most antimalarials), artemisinin acts like a less-discriminative “bomb” that detonates upon activation to cause widespread damage. The specificity of artemisinin may therefore be seen to be based on its activation rather than on its targets. These unique properties of artemisinin make it almost the ideal weapon against malaria, especially in combination with other drugs that act via distinct mechanisms and complement the pharmacological profile of artemisinin. An obvious advantage of a promiscuously targeting drug is also worth noting here: The development of drug resistance is much more difficult when mutation in one or a few specific targets is not sufficient to seriously impact drug activity. This advantage could well explain why artemisinin has remained generally efficacious despite its ubiquitous use over decades.

Nevertheless, recent trends have signaled the incidence and rise of malaria that is being cleared more slowly by ACTs, especially in the Asian endemic regions [69]. This topic has been comprehensively covered from various angles by recent reviews and commentaries [69–75]. Regardless of the controversies about the exact definition of “artemisinin resistance” in the field, the threat is undoubtedly real, given the place that artemisinin occupies in the control of malaria [76,77]. To resolve this burning issue, two major challenges must be overcome: 1) A full understanding of the MOA of artemisinin must be achieved; and 2) the genetic and physiological features of the newly emerged artemisinin-resistant strains must be defined. Even though the MOA of artemisinin has been largely demystified in the past few years, the molecular characterization of artemisinin-resistant malaria is far from clear. Continued efforts are required to achieve a complete picture of how artemisinin resistance relates to its mode of action. Based on this new knowledge, new therapeutic strategies can then be developed and tested.

4. Repurposing artemisinin

Artemisinin therapy is characterized by its outstanding tolerability and relative affordability. This combination of proven safety and accessibility make artemisinin a drug of exceptional interest for repurposing studies. Indeed, interest in non-malarial applications of artemisinin has increased steadily over time since artemisinin was first made known to the world [78]. While malaria remains the only disease for which artemisinin is an approved treatment, the potential applications of artemisinin in anti-cancer, anti-inflammatory, anti-parasitic (outside of malaria) and anti-viral roles, among others, have been explored in earnest over the years [78–82]. Here, we briefly comment on some promising research in artemisinin repurposing, especially in the field of cancer treatment, as a window into future drug development.

The efficacy of artemisinin in cancer cultures was first reported in 1993, and has since been expanded on and extensively characterized [83–85]. It is now well-reported that artemisinin and its derivatives display selective cytotoxicity against a range of cancer types in both in vitro and in vivo studies [86]. Forays into clinical testing have been generally promising, if limited in number and scale [87–89]. More than two decades of research on the basis of artemisinin action in cancer has uncovered a plethora of implicated targets and mechanisms. Artemisinin has been reported to induce mitochondrial apoptosis and other forms of cell death such as necroptosis, inhibit cancer angiogenesis and metastasis, and arrest the cancer cell cycle [90–97]. These outcomes are reportedly mediated by a combination of oxidative damage, DNA damage, alteration of gene expression, and interactions with a wide array of signaling pathways including mammalian target of rapamycin (mTOR), NF-κB, mitogen-activated protein (MAP) kinas, and Wnt/β-catenin, among many others [82,98–102]. These pathways and mechanisms have been extensively reviewed in recent publications [79–82].

While pathway validation is an important aspect of mechanistic study, it is also necessary to consider the big picture in terms of unifying drug activation and downstream activity in a manner similar to what was done in malaria studies. As is the case with malarial parasites, the activation mechanism of artemisinin in cancer cells is likely to be heavily linked to its specificity of action. Thus, the role of free ferrous iron versus free redox-active heme is once again being put under scrutiny, especially considering that iron is intimately linked to artemisinin-induced cytotoxicity in cancer [103,104]. Recent studies have once again shed light on
the role of heme in artemisinin activation in cancer cells, thereby drawing parallels with the case in malaria. In particular, a range of methodologies have been used to demonstrate that modulation of heme synthesis and availability clearly correlates with cytotoxicity [105–108]. It is also important to note that cancer cells have been reported to possess enhanced levels of heme metabolism and synthesis, and that this could underpin the cancer specificity of artemisinin in a similar manner to the case in malaria [109–111]. Specific targeting of artemisinin to mitochondria (the site of mammalian cell heme synthesis) or enhancement of heme levels by treatment with the heme precursor aminolevulinic acid (ALA) both improved anti-cancer activity [112–114]. A heme-centric mechanism of activation and an iron-dependent mechanism of downstream cytotoxicity could possibly be a point of reconciliation between the roles of those two species in the anti-cancer activity of artemisinin [115]. Further work to fully understand the basis of artemisinin specificity in cancer will be critical for future therapeutic applications.

At the same time, it is necessary to consider the appropriate direction when moving forward in terms of validating artemisinin MOAs in cancer. Consider the case in malaria, where artemisinin is proposed to indiscriminately attack adjacent targets upon activation. If artemisinin is activated in a similar manner in cancer cells, it is plausible that the same promiscuous multi-target mechanism would take place. This would explain the remarkable range of cellular effects and implicated pathways that have already been reported, as multiple targets and functional pathways are likely to be simultaneously affected by such a mechanism. Indeed, recent unbiased studies of artemisinin cancer targets using proteomics approaches have revealed a similar multi-target MOA by artemisinin in cancer cells [48,113,114]. The mechanism of cytotoxicity itself is also a matter of great interest, especially with regard to non-apoptotic forms of cell death. Recent work has closely linked artemisinin-induced cytotoxicity to oxidative damage and lysosomal function, with a focus on the role of iron in contributing to the iron-dependent form of cell death known as ferroptosis [116–118]. In particular, lysosome-mediated degradation of ferritin under autophagy conditions (termed ferritinophagy) releases free ferrous iron, which in turn contributes to both ferroptosis and iron-mediated generation of ROS [93,119]. Autophagy itself is a cellular process that is reportedly activated by artemisinin, but hasambiguous effects on cancer cell survival and the cytotoxicity of artemisinin [115,119]. It is clear that the relationship between autophagy, lysosomal activity, free ferrous iron, and iron-dependent ferroptotic cell death following artemisinin exposure represents a major area of uncertainty in the anti-cancer mechanism of artemisinin. However, efforts in unveiling novel, cancer-specific targets and mechanisms are steadily ongoing and continue to contribute to a grand view of artemisinin as an anti-cancer drug. Artemisinin-mediated effects on cancer stem cells, immunomodulation, cancer metastasis, cancer metabolism including the regulation of glycolysis, and a plethora of signaling pathways including signal transducer and activator of transcription 3 (STAT3), NF-κB, mTOR, and CREBP signaling are among recent reports, and indicate novel directions for further validation [115,120,121]. In particular, the potential ability of artemisinin to serve as an immunomodulator in cancer by regulating regulatory T cell (Treg) activity and the production of pro-cancer-survival immunosuppressive cytokines such as prostaglandin E2 (PGE2) is noteworthy, given the complex role of immunomodulatory drugs in cancer therapy [122–125]. Finally, efforts to improve the formulation and delivery of artemisinin-based drugs have shown promise in delivering enhanced efficacy and reduced susceptibility to drug resistance. These results include novel synthetic dimers, trimers, and drug conjugates (especially transferrin-conjugated systems), in addition to combination therapies; they represent an exciting ongoing area of research that has been reviewed comprehensively in recent publications [126–135].

In addition to the possible applications of artemisinin in cancer treatment, active research is taking place on its potential roles in addressing a range of other diseases. In particular, anti-inflammatory effects against autoimmune diseases and allergic asthma, among other conditions, have been reported in a range of disease models [78]. Some of these results correlate with observations of immunosuppression in patients undergoing artemisinin therapy for malaria [136]. Strong anti-viral effects of artemisinin have also been reported in herpes and in hepatitis B and C viruses, and other parasitic diseases including schistosomiasis have also been shown to respond to artemisinin treatment [137–141]. Recent findings have even identified a remarkable—if controversial—role of artemisinin in diabetes through inducing transdifferentiation of pancreatic β cells to generate γ cells [142,143]. The MOA for these alternative applications is frequently discussed in terms of the canonical model of ROS generation and oxidative damage induction upon endoperoxide cleavage; however, non-canonical (including endoperoxide-independent) mechanisms have also been proposed, especially in the case of immunomodulation [78,144]. It will be essential to pursue a clear view of how drug mechanisms and functions may differ under varying applications and conditions, while considering the importance of the conditions of drug activation. It is also worth noting that repurposing research might be best carried out in patients and regions that are not burdened with or at risk of malaria, in order to avoid possible interference or complications. Every care must be taken to ensure that the full potential of artemisinin can be realized without compromising its current applications.

5. Conclusion

The artemisinins are a class of remarkable drugs that have redefined the landscape of antimalarial therapy. A combination of outstanding potency, safety, and accessibility has put artemisinin at the forefront of the ongoing battle against the malaria scourge, where it has already impacted millions of lives. Since its discovery, a concerted effort by the global community has assembled a picture of a drug with a unique set of properties that makes it almost the ideal antimalarial drug. Active research in other fields has also revealed a broad spectrum of promising applications for artemisinin outside of malaria. We believe that it is only logical to seek to maximize the utility of this drug in a range of capacities. In the context of malaria, doing so means to continue to clarify the mechanisms of activation and action of artemisinin, while working to further improve its pharmacological properties both alone and in combination [145]. Combined with a firm grasp of the principles of artemisinin activity, this could be the key to clearing the uncertainties of artemisinin resistance. Such efforts would ensure that the drug can continue to perform in a similar or even greater capacity within the role that it has served for so long. Looking ahead, repurposing studies driven by a robust understanding of differential MOAs in different diseases and systems will also be instrumental in defining the future of artemisinin. Ultimately, it is our sincere hope that this gift from Chinese medicine can continue to serve the pursuit of health for people all around the world, for many years to come.

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