O

nly four months after China reported its first COVID-19 case to the World Health Organization (WHO) the virus had spread to every nation on the African continent. Despite being home to 17% of the world’s population, Africa currently accounts for just 2.5% of COVID-19 related deaths¹. But the pandemic may well have caused many more to die, not from coronavirus, but from malaria.

The 2020 World Malaria Report warned that disruptions to malaria prevention and treatment caused by the coronavirus could see as many as 100,000 malaria-related deaths in Sub-Saharan Africa². Similar effects were unfortunately experienced during the 2014-2015 outbreaks of Ebola in West Africa³. While most strategies being employed to control malaria have worked well, progress to reduce its incidence has stagnated. New strategies are needed to prevent the mortality rate from increasing further and to better prepare countries in the face of other unexpected pressures.

With the ambitious goal to create a world free of malaria, one not-for-profit research consortium—Target Malaria—is developing novel technologies using genetic modification to control the numbers of the malaria-transmitting mosquitoes.

THE NATURE OF MALARIA AND THE MICROBE RESPONSIBLE

Malaria is a disease that starts with a small single-celled parasite. This microorganism belongs to the genus *Plasmodium*, and of the four species that threaten humans, *P. falciparum* and *P. vivax* are the most common, and the former the most dangerous⁴.

Female mosquitoes alone spread malaria in nature. An infected mosquito injects a small number of parasites into its victim’s bloodstream while it feeds, and the parasites then travel to the liver where they multiply rapidly before infecting red blood cells. Flu-like symptoms begin when the parasites break out of the blood cells, one to four weeks after the bite.
Of the 229 million confirmed malaria cases worldwide in 2019, 94% occurred in Africa². Even more devastating, of the 409,000 malaria-associated deaths, 84% occurred in children under the age of five.

While there are more than 3500 species of mosquito worldwide and 837 in Africa, three very closely related species are responsible for most transmission of the disease: *Anopheles gambiae*, *Anopheles coluzzii*, and *Anopheles arabiensis*. These three species belong to the *Anopheles gambiae* complex which, if targeted, is likely to have the largest effect on the transmission of malaria. This species complex has tightly evolved with humans and is the key vector for malaria in sub-Saharan Africa.

Current vector control tools such as insecticides, bed nets, and drugs have been effective in reducing malaria cases but not in eradicating the disease. Target Malaria’s approach is meant to be complementary to the existing interventions by focusing on malaria control by mosquito control.

**RESEARCHERS UNIFY UNDER TARGET MALARIA**

In 2003, Prof. Austin Burt published a seminal paper⁵ describing the principle of genetically modifying a population of mosquitoes for applications in the control of vector-borne diseases. Prof. Burt predicted that malaria-transmitting mosquitoes could potentially disappear in an area within two years when using these novel genetic tools. It was fortuitous that only a few years prior in Imperial College London, where Prof. Burt was working, a team lead by Prof. Andrea Crisanti had created the first reliable system for germline transformation of a malaria-transmitting mosquito⁶.

The two groups were brought together in 2005 with a grant given as part of the Grand Challenges in Global Health initiative and in a little over a decade the team’s scientific progress had resulted in a new mechanism for genetic control measures within *An. gambiae*⁷.

The initial group of researchers has now grown into a team of 180 project members, with collaborating research partners in Africa, Europe, and North America. As the work progressed, the project grew under its new brand—Target Malaria.

Today, Target Malaria is working with five African partner sites: Burkina Faso, Cape Verde, Ghana, Mali, and Uganda. This work is being headed by Dr. Abdoulaye Diabate at the Institut de Recherche en Sciences de la Santé (Research Institute on Health Sciences), Bobo-Dioulasso, Burkina Faso; Dr. Adilson de Pina at the Instituto Nacional da Saúde (National Institute of Public Health, CCS-SIDA), Cape Verde; Dr. Fred Aboagye-Antwi at the University of Ghana, Accra, Ghana; Dr. Mamadou Coulibaly at the University of Bamako Malaria Research and Training Center, Bamako, Mali; and Dr. Jonathan Kayondo at the Uganda Virus Research Institute, Entebbe, Uganda.
Each of the African partners and their teams bring a different skillset to the collaboration, none being able to deliver all key elements independently. A relationship of co-development is fostered within the project with scientists working together across countries and with communities towards a common objective and vision—a world free of malaria.

The three key guiding pillars

Target Malaria’s work is structured around three key pillars: science, stakeholder engagement, and regulatory affairs. Each pillar is essential for the project’s success, supporting responsible research and development of genetic technologies, a commitment to engaging a wide variety of stakeholders, as well as ensuring compliance with all national regulations and laws.

SCIENCE

To date, Target Malaria has made significant scientific advancements on the path to developing a new tool for vector control for malaria. Researchers have demonstrated the proof of concept in creation of a transgenic sterile male *An. gambiae* strain, demonstration of the ability to modify a laboratory population of *An. gambiae* mosquitoes to be male biased⁹, suppression of a small cage population of laboratory reared *An. gambiae* mosquitoes¹⁰, creation of the first gene drive mosquitoes capable of suppressing a laboratory population of *An. gambiae* mosquitoes¹¹, modelling the potential of genetic control of malaria mosquitoes¹², modelling suppression of malaria vector using gene drive¹³, and the importation of the first genetically modified mosquitoes (a self-limiting sterile male line) into Burkina Faso for contained laboratory use in 2016 and regulatory approval for the subsequent release of the same self-limiting sterile male line in 2019.

STAKEHOLDER ENGAGEMENT

The list of stakeholders is vast, from grass roots, those local communities where the project is working, through to local civil society organizations, regional governing bodies, and the appropriate governmental agencies, all in-country, as well as a range of interested parties outside of the African partner countries. Target Malaria is committed to ensuring that the stakeholders understand the research and long-term goals of the project enabling them to make an informed decision on whether to support the project’s efforts. Engagement also helps ensure that the research is welcomed and useful in the fight against malaria. Most importantly, Target Malaria will learn a lot from their stakeholders through the process.

REGULATORY AFFAIRS

While an important aspect of Target Malaria’s strategy is to focus on the communities that might benefit from the technology and that are most concerned by the research activities, they also have an ongoing, transparent dialogue with other stakeholders at the national and international level. For example, the project is taking a phased approach to its development pathway in line with guidance from the WHO¹⁴.

TECHNOLOGY BEING DEVELOPED

The goal is to develop modified mosquitoes that can pass on to their progeny a self-sustaining genetic change, a process aiming to reduce specific mosquito populations to break the malaria transmission cycle. To do so, Target Malaria is using gene drive, a phenomenon that occurs in nature and causes a selected trait to rapidly increase in frequency through a population via sexual reproduction over several generations. Gene drive works by increasing the likelihood—from the usual 50 per cent to greater than 95 per cent—that a modified gene will be inherited by its offspring. This means that over the course of several generations, a selected trait could become increasingly common within a specific species (depending on the specific area and how the animals move around within it).

Researchers are investigating the use of genes that produce enzymes that cut specific sequences of DNA. Called nucleases, these enzymes found in simple single celled organisms can copy themselves from one chromosome to another⁸. When introduced into the malaria mosquito, the nuclease works by identifying and cutting a selected site within essential genes targeted by researchers, rendering them functionless, such as reproductive genes. The subsequent effects depend on the nature and importance of the gene.

Target Malaria’s goal is to produce modified malaria mosquitoes that can pass these genes on to greater than 95 per cent of their offspring, so the modification is spread throughout the specific population relatively quickly and is effectively “self-sustaining”. This strategy is known as population suppression, and as the mosquitoes themselves do the work of spreading the modification, it makes the reduction of the malaria mosquito population relatively cost effective and simple to implement.

Current gene drive research is at an early stage, and so definitive decisions about gene drive-based tools are premature. Based on current progress, field releases of a gene drive-based tool are many years away. This gives scientists and stakeholders, specifically those from countries where gene drives might one day be employed, valuable
time to consider the important questions of regulation, risk assessment, ethics, and engagement, and to prepare for assessing any application related to gene drive mosquitoes and their potential use as a tool for vector control for malaria.
AN ECOLOGICAL APPROACH SPEARHEADED IN GHANA

As the gene drive approach in development by Target Malaria will specifically target the *An. gambiae* complex to reduce its population, it is vital to ensure there are no undesirable consequences to the rest of the plant and animal communities. In Ghana, researchers are focusing on the ecological implications of the work; the role of the *An. gambiae* mosquito in the broader ecosystem. The ongoing research in Ghana aims to predict these potential effects.

While some aspects of *An. gambiae* ecology is well studied, research in Ghana will provide a more complete picture, specifically determining the interactions between *An. gambiae* and other mosquito species as well as predators, prey, and vertebrate hosts. In this sense, the research is based on a community ecology approach rather than looking at just mosquito ecology.

Researchers are sampling, as far as they are able, insects from the entire aerial communities across all habitats, not just where they expect to find high numbers of *An. gambiae*. Equally true for insectivores that might feed on mosquitoes or similar small aerial insects, they are taking fecal samples or stomach contents. In aquatic habitats, they are sampling from a range of water bodies and collecting representatives across all insect and insectivorous groups. And, because adult mosquitoes rely on flower nectar for food, they are sampling the pollinator community as well. Finally, they are looking at the community of biting flies and their vertebrate hosts by determining blood-meal interactions to better understand shared hosts and the potential for zoonotic disease transmission\(^{15}\). Importantly, all methods and target sample numbers have been cleared by independent ethics boards at both the University of Ghana and the University of Oxford to ensure there is no lasting impact on the community of plants and animals where the work takes place.

Using this broad community approach allows researchers not only to describe the role of *An. gambiae* in the ecosystem but also to predict how the rest of the ecological community would respond to *An. gambiae* reduction. For example, the data would allow them to determine which insects might face more pressure from predators if those that feed on mosquitoes shifted their feeding behaviour to replace *An. gambiae* in their diet and how this change might affect the rest of the food web. As there is no known animal or plant that relies solely on *An. gambiae*\(^{16}\), and food webs tend to rewire following minor perturbations\(^{17,18}\), it is predicted that there will not be any significant effects because of *An. gambiae* population reduction. Regardless, it is necessary to ensure this is the case and a community ecology approach will make this possible.

Building a DNA barcode reference library for Ghana

The tools being used to construct the food web are mostly molecular-based, all requiring the creation of a DNA barcode library for insects in the area as the important first step. To this end, researchers in Ghana regularly collected terrestrial insects from villages in the southeast of Ghana for a year.

Once the library is established, they can then start looking at the feces and the gut contents of insectivores
and use that library to match and identify prey DNA fragments using DNA metabarcoding. Researchers are using a similar metabarcoding approach for the aquatic food web while for the pollination network they will use a combination of traditional observational methods and DNA metabarcoding of pollen from caught insects. Finally, for the blood meal analysis, they are metabarcoding the blood meals of fed mosquitoes and other biting flies to identify what has been bitten.

The ecological research in Ghana is a collaboration between Dr. Fred Aboagye-Antwi at the University of Ghana, Prof. Sir Charles Godfray and Prof. Owen Lewis at the University of Oxford, as well as an extensive team of postdoctoral researchers, Ph.D. students, and technicians at both institutions. Part of the UK team, postdoctoral researcher and project coordinator, Dr. Talya D. Hackett is organizing efforts between countries, including a collaboration with the Centre for Biodiversity Genomics (CBG) in Guelph, Canada, global leader in the field of DNA barcoding. Supported by Prof. Paul D. N. Hebert, the Director of CBG, and Dr. Michelle L. D’Souza, samples from Ghana have begun making their way to the large sequencing platforms housed at the CBG.

So far, about 3000 insects have been processed and 530 BINs (species proxies) have been documented, about 70% of which are unique to the project. Efforts will ultimately barcode 100,000 specimens and fill a large gap in barcode data currently missing from West Africa.

**CONCLUSIONS**

Examining diets to determine species-specific interactions in a complex community food web is only possible at this large scale with molecular techniques, and only recently, because the costs of DNA barcoding and metabarcoding techniques have dropped. Even five years ago this sort of a project would not have been feasible.

Apart from building large, comprehensive food webs, these data can further inform our understanding of things like community structure and insect population dynamics, the dietary overlap of insectivorous species, and niche overlap of different mosquito species.

All data will be made publicly available. Ultimately, this project is creating a wealth of information, not just for Target Malaria’s research goals, but for the broader scientific community and for other people within Ghana and West Africa.

These efforts are a demonstration of the power of DNA barcoding and its ability to reveal the nature and intensity of interactions among all species. This endeavour, to reveal species interactions to clarify their role in structuring biological communities, is a key research theme of BIOSCAN, iBOL’s new seven-year, $180 million global research program that aims to revolutionize our understanding of biodiversity and our capacity to manage it.

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_Images show some of the faces of the research team in Ghana: Dr. Fred Aboagye-Antwi (Ghanaian principle investigator, top left), Helen Selorm Wohoyie (Assistant stakeholder engagement and communications advisor, top center), Divine Dzokoto (Senior stakeholder engagement and communications advisor, top right) and Dr. Talya D. Hackett (project coordinator), Bernard Aye Adams, Ezekiel Yaw Donkor, and Naa Na Afua Acquaah (laboratory technicians) (bottom, left to right). Photo credit: Lema Concepts Africa_
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