The reintroduction of the mosquito *Aedes aegypti* Linn. in Brazil in the 80’s started a period of dissemination of diseases transmitted by this mosquito. Upon every new serotype or disease introduced in this country, new epidemics arise. Recently, two new diseases transmitted by this mosquito were introduced in Brazil, namely chikungunya and Zika. Especially, Zika has alarmed health authorities worldwide due to its sexually-transmitting capabilities, microcephaly in newly born babies, and the Guillain-Barré syndrome. These data show that there is an urge to deepen the research not only for treating these diseases, but also to control its mosquito vector. Considering the resistance that the larvae of *A. aegypti* have developed to the larvicides currently used, the development of studies aimed at obtaining effective compounds against the larvae of this mosquito vector are of fundamental importance for human society. In this context, our research group synthesized *N*-tosylindole, a molecule with sufficient larvicidal potency to be directed to
pharmacotechnical development studies. To carry out the studies that are being proposed, this project initially aims to carry out the synthesis on a larger scale of the compound in question, followed by the development of a formulation for its use, analyzes of its cytotoxic and genotoxic potential, as well as its residual effect. Cellular and genetic toxicity analyzes will be carried out in HepG2/C3A human hepatoma cells, with extension to analyzes of effects on the cell cycle, apoptosis and expression of genes involved in these metabolic pathways. Nanotechnology will be used to obtain a formulation, aiming to increase the effectiveness of vector control and enable the use of a molecule with low water solubility, N-tosylindole, making this product viable for use in water. A suitable formulation must release the larvicide continuously in order to maintain the concentration of the larvicide at a high enough level to maintain its effect for a long time. The residual effect will be obtained in laboratory aiming to determine the number of days its active ingredient stays in the environment in which it was placed. This same study can be increased to a semi-field scale, evaluating the residual effect in conditions more similar to reality in semi-field evaluation using containers in the environment. Then the formulation will be evaluated for impacts on non-target plant species via evaluation of N-tosylindole effects on germination and plant growth. At the end of the project, it is expected that the data collected will allow the use of the molecule in question as a larvicide and serve as another tool in the control of the A. aegypti mosquito and in the fight against the diseases vectored by it.

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Data Collection

What data will you collect or create?

Data will be obtained from three experiments listed below:

1. Formulations containing N-tosylindole characterized by polarized light microscopy, small angle x ray scattering, rheological analysis and Fourier-transformed infrared spectroscopy;
2. Germination and growth of seeds exposed to different concentrations of the title compound. Number of germinated seeds, size and weight of shoots;
3. Residual effect in laboratory and field controlled situations. Length in days the test compound produces larvicidal effect in a container with water. Number of dead larvae with time.
4. Toxicity data will be obtained from the cultured lymphocytes and HepG2/C3A human cells treated with different concentrations of N-tosylindol. The data will be shown the effects of these compounds in the context of cytotoxicity, genotoxicity, (with active liver metabolizing enzymes) and human leukocytes, and the effects of these compounds on the progression of the cell cycle, induction of apoptosis, and gene expression in HepG2/C3A cells (human hepatoma).

All data collected will be in tables and laboratory chains.

How will the data be collected or created?

Formulation data will be collected using equipment mentioned before, seed germination and growth data will be collected by germinating seeds in petri dishes in controlled environment, residual effect data will be collected using beakers (lab setting) or containers (semi-field setting) containing water, formulation, and frequently added *Ae. aegypti* larvae. The data about toxicity will be collected from cultured human cells (lymphocytes and HepG2/C3A) exposed to N-tosylindol.

Documentation and Metadata

What documentation and metadata will accompany the data?

Graphics, FIDs, spectra, figures, and tables.

Ethics and Legal Compliance
How will you manage any ethical issues?

All procedures involving toxicity analysis will be approved by the Human Ethics Committee of the Faculty of Philosophy and Sciences, UNESP, Marília town.

Insect rearing and larvae production procedures were approved by the Ethics Committee of Animal of Federal University of Sergipe (CEPA-UFS) application # 4386260321.

How will you manage copyright and Intellectual Property Rights (IP/IPR) issues?

As a result of this project, articles will be published. Copyright will be determined jointly with the members of our research group.

Storage and Backup

How will the data be stored and backed up during the research?

Data will be backed up with copies in hard drives in laboratory computers, as well as the university offers google Drive, which assists in data storage.

How will you manage access and security?

All researchers involved in the project will have access to the data, but only the researcher responsible and coordinator will edit and update the backup.

Selection and Preservation

Which data are of long-term value and should be retained, shared, and/or preserved?

The data will be shared with the scientific community through publications. In addition, they will serve to originate future projects.

What is the long-term preservation plan for the dataset?

We store these data in drives of all laboratories involved in this project, and laboratory chains.

Data Sharing
How will you share the data?

The data will be used for participation in a scientific congress. And, after a great understanding of the results, the data will be published in the form of an article or patents.

Are any restrictions on data sharing required?

Before intellectual protection the data is not available. After intellectual protection the data cannot be used for commercial purposes without consent from the holder. Toxicity data will be published for the entire community, after which it can be shared.

Responsibilities and Resources

Who will be responsible for data management?

The data acquired during the project will be managed by the responsible researcher in each laboratory and the project coordinator.

What resources will you require to deliver your plan?

The project will be necessary for collaboration with other researchers to collect and analyze the data obtained from the mentioned experiments.