## RTS,S MEDICAL AND SCIENTIFIC QUESTIONS & ANSWERS

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## **Disclaimer**

The primary focus of this resource is to be an internal training tool for RTS,S malaria vaccine candidate, containing related data in the format of a Q&A for Medical Affairs personnel. Information presented here is not for external distribution.

Whilst this document can be inspirational for reactive responses to experts or medical enquiries, local regulations, the GSK Code of Practice, scientific engagement principles and/or medical information processes should be followed appropriately.

## **Please Note**

- For media enquiries, please refer to the specific reactive Q&A for Media Enquiries and notify the Global Pipeline Communications team before you respond to a request for an interview so that they can help you to prepare (contact person: Aoife Pauley at aoife.x.pauley@gsk.com).
- The vaccine RTS,S/AS01 has completed phase 3 clinical program and positive regulatory assessment from the European Medicines Agency, but is not yet authorized for marketing in any country. The RTS,S vaccine is being developed in Public Private Partnership with PATH-MVI, as an additional tool to be added to the currently available malaria preventive interventions and for implementation through the national immunization programs in malaria endemic regions in sub-Saharan African countries.
- When referencing clinical data on RTS,S any statements should be prefaced by "In this study...", to make it clear that it is too early to make any general statement on the vaccine profile outside the context of the ongoing clinical trails.
- Have you found what you were looking for? If you have any suggestions for information which should be included in this tool please contact us at the following address: Carys Calvert at <u>carys.calvert@gsk.com</u>.

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Does the use of adjuvants in the RTS,S malaria vaccine candidate increase the risks or side effects associated with the vaccine?

Adjuvant Systems have been used effectively in the development of the RTS,S malaria vaccine candidate for more than 15 years. The Adjuvant Systems used in RTS,S development have demonstrated a safety profile comparable to that observed with other vaccines routinely given to infants through the well-established Expanded Programme for Immunisation. GSK will continue to assess the safety and efficacy of the adjuvanted RTS,S malaria vaccine candidate as part of the ongoing clinical trials, as well as during post-approval pharmacovigilance studies.