

WHO Prequalification of Vector Control Products

Annex V. Spatial emanator studies

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Table 1.2 Methods

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Free-flight room

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Table 1.3. Results

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Table 1.1 General		
Report section	Description	Critical parameters to report
	General	
Cover page		
Table of contents		
GLP compliance statement	An official statement of compliance with GLP requirements. The GLP certificate can be provided as part of this section or as an annex to the report	
Results summary	Briefly summarise the results and conclusions of the study. This can be in tabular or narrative text format.	
List of abbreviations	List of abbreviations used in the study report. The use of abbreviations should be kept to a minimum.	
Background	Relevant background information for the study. This can be a brief description of the product and its proposed use.	
Study rationale	A brief description of the rationale for conducting the study and the intent of its use	
Study objectives	List the objectives of the study. Study objectives should be clearly written and described. If the study has been conducted to meet the requirements of multiple bodies, the full list of study objectives can be provided in this section, with those study objectives related to the prequalification product assessment clearly indicated.	
Study endpoints	This section should list and describe all endpoints used in the study, including descriptions of primary and secondary endpoints where relevant.	Primary endpointsSecondary endpoints
	If multiple strains of test systems have been tested in the study, identify the	



Table 1.1 General		
Report section	Description	Critical parameters to report
	General	
	test system which was used to determine the validity of the study/provide the scientific determination of product performance, and provide a rationale for the selection of said test system as the decision-making strain. Endpoints should be used consistently throughout all data generation for a product, with the exception of early exploratory studies which might be submitted in a dossier as supplementary evidence.	
Criteria for study acceptance	 List and describe the criteria for Acceptance of the study as scientifically valid Evaluation of the product as having met the requirements for prequalification for that particular study type 	 Criteria for controls Criteria for evaluation of the proposed product as having met the requirements for prequalification for that particular study type, e.g. laboratory assessment
Guidance and protocol deviations	Provide any deviations from either the study protocol (as per GLP requirements) and/or from WHO guidance	 Deviations from the study protocol As per GLP facility requirements Deviations from WHO guidance Evidence-based justifications/rationales Assessment of the impact on study validity, acceptability, robustness, with additional evidence to support the assessment where necessary Any adjustments that were made to the study protocol in response to considerations received from WHO as part of a protocol review submission



Report section	Description	Critical parameters to report
		Methods
1.2.1 Test systems	:	
Test systems	Description of the test systems used in the study	 Colony maintenance and brief summarised rearing procedures Light cycle of insectary Age and physiological status used in bioassays If multiple bioassays have been used, report the age and physiological status for each method separately Most recent date of insecticide resistance characterisation (note that the results of the characterisation should be reported in the matrix for mosquito strains) Justification for the selection of test system(s), including reference to the product AI and mode of action, and the characteristics of the test system(s) that make it a suitable choice
1.2.2 Study sites		
Description and selection of study sites (for semi-field studies)	Narrative description of semi-field study site(s), including a justification for the site(s) suitability	 Location GPS coordinates Description of seasonal variations and rainfall
1.2.3 Characterisa	tion of vector population(s)	
Characterisation of local vector population (for semi-field studies)	Description and characterisation of the local vector population at semifield sites, including suitability for use in testing the proposed product	 Vector species and composition, including sibling species if present Description of insecticide resistance status and mechanisms (note that the results of the characterisation should be reported in the matrix for mosquito strains)
1.2.4 Test items, p	roduct information	
Test and reference items	Description of the batch(es) of test and reference items used in the study.	 The number of batches of test items used in the study All batch numbers for test and reference items The number of test and reference items received at the testing facility The number of test items received per batch of test items Source of all test and reference items



Report section	Description	Critical parameters to report
		Methods
Test and reference items	Description of the product composition and AI(s) of test and reference items	 Date of manufacture Date of receipt at the testing facility Storage conditions post-receipt Justification for the choice of positive control(s) Material Product type, e.g. active vs passive emanation Al description Name Mode of action Dosage
1.2.5 Sample prep	paration	
Sample preparation	Description of sample preparation	 The number of product samples used in the study Sample size(s) Sample storage conditions Description of product ageing, e.g. natural/artificial/operational Environmental conditions (temperature, relative humidity, airflow, air exchange) Storage conditions Sample storage and shipment details for chemical analysis
1.2.6 Insecticide r	esistance status	
Insecticide resistance status of test systems and local vector populations	If insecticide resistance characterisation of test systems has been conducted as part of the study, describe the method. For semi-field studies, describe the methods used to determine the LC ₅₀ and LC ₉₀ to the AI(s) used in the proposed product.	 Insecticides tested Insecticide dosages Method used, i.e. WHO tube test or bottle bioassay Total number of mosquitoes tested Number of mosquitoes per replicate Number of mosquitoes per test arm Exposure duration Post-exposure holding conditions and monitoring Method for determining LC₅₀ and LC₉₀ (for local vector populations used in semi-field studies)



Report section	Description	Critical parameters to report		
Methods				
Bioassay methods	Description of bioassay method used: Free-flight rooms	 Free-flight chamber design and measurements Environmental conditions (temperature, relative humidity, airflow, air exchange) Starvation protocol Number of products per room and installation location(s), e.g. height, distance from walls, etc. Acclimatisation time and temperature(s) for test systems and materials Time of day when tests were conducted, including start and end times Exposure duration Product storage conditions between test days Post-exposure holding duration and environmental conditions in holding room Endpoint recording Holding receptacle Timing and placement of sugar sources 		
Bioassay methods	Description of bioassay method used: Experimental hut	 Hut design and measurements Study arms Decontamination and/or refurbishment procedure Baseline collections and scavenging rate estimation method Number of products per hut and installation location(s), e.g. height, distance from walls, etc. Product storage conditions between testing nights Method for recording environmental conditions (temperature, relative humidity) Acclimatisation time for materials Time of day when tests were conducted, including start and end times Exposure duration Transport protocol for collected mosquitoes Mosquito identification procedure Post-exposure holding duration and environmental conditions in holding room Endpoint recording Holding receptacle 		



Report section	Description	Critical parameters to report
		Methods
		Density of mosquitoes in holding receptacleTiming and placement of sugar sources
1.2.8 Study design	1	
Study design	Free-flight room studies	 Ethical review board permission Number, gender and age range of volunteers Collection period Cleaning protocol Number of mosquitoes per replicate Total number of replicates and number of replicates per test arm Mosquito collection method Mosquito scoring method Endpoint recording Adverse effects monitoring (if indicated)
Study design	Semi-field studies	 Ethical review board permission Environmental conditions (temperature, relative humidity) Number, gender and age range of volunteers Latin square design Treatment allocation Volunteer allocation Volunteer rotation Collection period Cleaning protocol Total number of collection nights Exposure duration Mosquito collection method Mosquito scoring method Endpoint recording Adverse effects monitoring (if indicated)
1.2.9 Sample size	calculations	
Sample size calculation for free-flight rooms	Provide a full description of the calculations employed to arrive at the required sample size(s)	 Data source used to parameterize sample size calculations, e.g. previous studies, simulated data Endpoint used to power study Point estimate used



Report section	Description	Critical parameters to report
		Methods
		 Procedure used to estimate the sample size, e.g. simulations, existing software/packages Details of the procedure that was followed Assumptions considered, e.g. effect size, power, variability, significance level, and justification(s) for the values of each assumption
Sample size calculations for semi-field studies	Provide a full description of the calculations employed to arrive at the required sample size(s)	 Data source used to parameterize sample size calculations, e.g. previous studies, simulated data Endpoint used to power study Point estimate used Simulation procedure used to estimate the sample size/number of required nights of collection Details of the procedure that was followed Assumptions considered, e.g. effect size, power, variability (e.g. differences between huts/chambers, sleepers, collection nights), significance level, and justification(s) for the values of each assumption
1.2.10 Data analy	sis	
Data analysis for descriptive statistical analyses	Description of the descriptive statistical methods used to summarise and describe data in the report, including measurements of dispersion	 Number of samples Number of mosquitoes per study arm Mean/Median (as appropriate) Standard deviation Range
Data analysis for inferential statistical analyses	Description of the fitted inferential model used for each endpoint (including secondary endpoints)	 For each endpoint: Type of model Type of endpoint/data Distribution Fixed effects (including the type of variable, e.g. continuous or categorial/factor Random effects Justifications for any deviations from published guidance



Table 1.3. Results					
Report section	Description	Critical parameters to report			
	Results	5			
Characterisation of local vector population(s)		 Composition of local vector population, including sibling species and seasonal variation (where appropriate) 			
Free-flight room studies	Narrative, tabular and graphical presentation of results of free-flight room studies	 Evaluation of the results in terms of compliance with the required sample size Summary results for all primary and secondary endpoints, presented by study arm with appropriate measures of dispersion (tabular) Inferential statistical results (tabular) Graphical presentation of results Narrative description of results The code used for statistical analyses in the format that it was produced (separate file(s)) 			
Semi-field studies	Narrative, tabular and graphical presentation of results of semi-field studies	 Evaluation of the results in terms of compliance with the required sample size Summary results for all primary and secondary endpoints, presented by study arm with appropriate measures of dispersion (tabular) Inferential statistical results (tabular) LC₅₀ of the local vector population to the AI(s) LC₉₀ of the local vector population to the AI(s) Graphical presentation of results Narrative description of results The code used for statistical analyses in the format that it was produced (separate file(s)) 			



Report section	Description	Critical parameters to report
	Discussion and co	nclusions
Discussion	For each study, an interpretative discussion of the results must be provided.	 Interpretation of the study results with reference to the criteria for study acceptability identified in Criteria for study acceptance, e.g. evaluation of the scientific validity of the study based on the parameters of the study and the results of controls Specific discussions on any methodological deviations, anomalies in results, or other factors which may have impacted the results should be included. Interpretation of the study/sub-study results with reference to the criteria for study acceptability identified in the Criteria for study acceptance with regards to the evaluation of the proposed product as having met the requirements for prequalification for that particular study type with specific discussions on any methodological deviations, anomalies in results, or other factors which may have impacted the results should be included.