FRPath.org Where the Roads to Accelerated Assessments Converge



FRPath.org Country and FRP Information Input Form				
Country: Suriname Agency Name: Ministerie van Volksgezondheid (Ministry of Health)				
Name of FRP: Click here to enter text.				
Is this FRP Proposed or Active? Choose an item.				
Date FRP was officially enacted: Click here to enter a date.				
1. Facilitates activities	2. Accel	erates the regulatory	3. Relies on or recognizes a prior	
during development	r	eview process	regulatory decision	
Is a Guidance or SOP describing how		Choose an item.		
to apply this FRP publicly available?				
When should the FRP be requested?		Choose an item.		
Does the agency provide		Choose an item.		
assistance/advice to the sponsor?				
For which types of product(s) can this		Click here to enter text.		
FRP be used? E.g. NMEs, generics,				
biologics, biosimilars, all products				
Must the product address an unmet		Choose an item.		
medical need or serious condition?				
If a fee is required, what is the		Click here to enter text.		
amount (in US\$ equivalent)				
Total target (agency) time for		Click here to enter text.		
assessment (calendar days)				
Total target (company) time for		Click here to enter text.		
responses to agency questions (If				
stated)				
Select one of the following (* see definitions at end of document)				
Is this a verification review		an abridged* review	Is this a full* review of all parts of	
(a recognition pathway)?*		ed dossier portions)?	the dossier?	
	(a re	liance pathway)?*		
			_	
If this is a reliance or recognit		☐ Click here to enter text	_	
pathway, what are the accep-		Click here to enter text	_	
pathway, what are the accept reference agencies?			-	
pathway, what are the accept reference agencies? How many reference agency		Click here to enter text	-	
pathway, what are the accept reference agencies? How many reference agency decisions are required?	ted		-	
pathway, what are the accept reference agencies? How many reference agency decisions are required? Does this FRP require submis	sion of	Click here to enter text	-	
pathway, what are the accept reference agencies? How many reference agency decisions are required?	sion of	Click here to enter text	-	
pathway, what are the accept reference agencies? How many reference agency decisions are required? Does this FRP require submis Assessment Reports from price	sion of	Click here to enter text	-	
pathway, what are the accept reference agencies? How many reference agency decisions are required? Does this FRP require submis Assessment Reports from prid decisions?	sion of or	Click here to enter text Choose an item.	-	
pathway, what are the accept reference agencies? How many reference agency decisions are required? Does this FRP require submis Assessment Reports from prid decisions? Is a CPP (Certificate of	sion of or	Click here to enter text Choose an item.	-	

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documentation to the CPP be used? If		
so, what types of documents?		
If this process is through a Regional	Click here to enter text.	
Regulatory Initiative, which countries		
participate in this process?		
Does the product have to have been	Click here to enter text.	
marketed in another country? For a		
specific amount of time? If so, for		
how long?		
How are queries to the companies	Choose an item.	
sent?		
Are external reviewers (e.g. non-	Choose an item.	
agency) involved in the assessment?		
Post-authorization study	Choose an item.	
commitments		
For how long is the initial approval or	Choose an item.	
designation valid?		
Any other details you wish to	Click here to enter text.	
provide?		
Date of this update	22 MARCH 2020	
References	1. <u>Suriname: Pharmaceutical Country Profile</u> .	

*Definitions:

Verification review: A checklist review based on recognition of a prior regulatory decision. Recognition is the routine acceptance of the regulatory decision of another regulator or other trusted institution. Recognition indicates that evidence of conformity with the regulatory requirements of economy A is sufficient to meet the regulatory requirements of economy B.

Abridged review: An abbreviated review of selected portions of the dossier and the reliance on prior assessment decisions. Reliance is the act whereby a regulatory authority in one jurisdiction may take into account/give significant weight to work performed by another regulator or other trusted institution in reaching its own decision

Full review: A comprehensive review of all components of the dossier. This may or may not be CPP-dependent. This may form part of a reliance or recognition pathway.

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