## FRPath.org Where the Roads to Accelerated Assessments Converge



FRPath.org Country and FRP Information Input Form					
Country: Chile Age					
Name of FRP: Priority Review: 9	Name of FRP: Priority Review: Short Registration Procedure				
Is this FRP Proposed or Active? Active					
Date FRP was officially enacted	Click here to enter a date.				
1. Facilitates activities during development	2. Accelerates the regulatory review process 3. Relies on or recognizes prior regulatory decision				
Is a Guidance or SOP describing how to apply this FRP publicly available?	Yes- see reference below				
When should the FRP be requested?	Choose an item.				
Does the agency provide assistance/advice to the sponsor?	Yes- For any product type				
For which types of product(s) can this FRP be used? E.g. NMEs, generics, biologics, biosimilars, all products	<ul> <li>Priority review is applicable for medicines for the following cases:</li> <li>For serious diseases and life threatening conditions, and when medicines are apparently expected to contribute to the improvement of quality of healthcare based on overall evaluation of the seriousness of the target disease and medical usefulness.</li> <li>When application is submitted for marketing approval and consideration is made based on the opinions of the external experts</li> </ul>				
Must the product address an unmet medical need or serious condition?	Yes				
If a fee is required, what is the amount (in US\$ equivalent)	Click here to enter text.				
Total target (agency) time for assessment (calendar days)	Click here to enter text.				
Total target (company) time for responses to agency questions (If stated)	Click here to enter text.				
Select one of the following (* see definitions at end of document)					
Is this a verification review (a recognition pathway)?*	Is this an abridged* review (selected dossier portions)? (a reliance pathway)?*	Is this a full* review of all parts of the dossier?			
		$\boxtimes$			
If this is a reliance or recognition pathway, what are the accepted reference agencies?	Click here to enter text.				

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How many reference agency	Click here to enter text.		
decisions are required?			
Does this FRP require	Choose an item.		
submission of Assessment			
Reports from prior decisions?			
Is a CPP (Certificate of	Yes at time of submission		
Pharmaceutical Product)			
required for approval?			
Can an alternate form of	Certificate is for those products imported during any stage of the		
reference documentation to	production process, issued by the corresponding health authority		
the CPP be used? If so, what	of the country where the manufacturing facilities are located		
types of documents?	certifying that the foreign manufacturer is duly authorized in their		
	country and follows Good Manufacturing Practices in accordance		
	with WHO recommendations. It also indicates that production		
	areas or types of products are authorized for manufacturing,		
	unless such information is included in the document such as		
	marketing authorization certificate, pharmaceutical product		
	certificate, marketing authorization certificate or official		
	certification recommended by the World Health Organization		
	issued by the authorities of the country of origin and duly		
	legalized.		
If this process is through a	No, this process is not through a Regional Regulatory Initiative.		
Regional Regulatory			
Initiative, which countries			
participate in this process?			
Does the product have to have	Click here to enter text.		
been marketed in another			
country? For a specific amount			
of time? If so, for how long?			
How are queries to the	Choose an item.		
companies sent?			
Are external reviewers (e.g.	Yes- always		
non-agency) involved in the			
assessment?			
Post-authorization study	Always required		
commitments			
For how long is the initial	4-5 years		
approval or designation valid?			
Any other details you wish to	- Any pharmaceutical product, whether imported or		
provide?	manufactured in the country, requires a sanitary		
	registration (marketing authorization) in order to be		
	distributed or used under any title in Chile (a		
	pharmaceutical product may be exceptionally authorized		
	by the ISP to be used temporarily without prior sanitary		
	registration if an epidemic, emergency or catastrophe		
	occurs, or if required for an urgent medical use or for		

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		scientific research or clinical trials).	
	-	In general terms, for the sanitary registration of a	
		pharmaceutical product the applicant will be required to	
		comply with general requirements including the	
		submission of administrative information, technical	
		information, pharmaceutical quality information and data	
		on safety and efficacy of the product. Special	
		requirements will also be applicable for fixed dose	
		combination products, pharmaceutical combination	
		products, phytopharmaceutical products; homeopathic	
		products and biologicals.	
	_	Sanitary registrations for pharmaceutical products will last	
		for a period of 5 years, which is renewable for equal and	
		successive periods, if not cancelled.	
	_	The marketing authorization of generic drugs can be	
		according to the ordinary approval procedure ("New Drug	
		Application"), the simplified approval process ("Simplified	
		Registration Procedure"), or the abbreviated approval process ("Short Registration Procedure"). The registration	
		procedure will depend on the pharmaceutical product	
		(NDA for new active ingredients, new doses or new routes	
		of administration; Simplified procedure if there are any	
		products already approved with the same ingredients in	
		the same dose and with the same route of administration;	
		and Short procedure if the Ministry of Health requests it	
		either for the implementation of public health plans and	
		programmes, or if the active ingredient is added to the	
		National Formulary).	
Date of this update	22 Febr	ruary 2020	
References	1.	Drug Approval System of Chile.	
		https://www.google.com/url?sa=t&rct=j&q=&esrc=s&sour	
		<pre>ce=web&amp;cd=1&amp;cad=rja&amp;uact=8&amp;ved=2ahUKEwidoefV60</pre>	
		PnAhUEahQKHS-	
		8CbwQFjAAegQIAhAB&url=http%3A%2F%2Fwww.nifds.	
		go.kr%2Fbrd%2Fm_95%2Fdown.do%3Fbrd_id%3Dboard	
		_mfds_411%26seq%3D22991%26data_tp%3DA%26file_s	
		<u>eq%3D1&amp;usg=AOvVawoFnA8qPnPrraG_uAZVAvbt</u>	
		Accessed on 22 February 2020.	
	2.	Regulatory, Pricing and Reimbursement.	
		https://pharmaboardroom.com/legal-articles/regulatory-	
		pricing-and-reimbursement-chile/ Accessed on 22	
		February 2020.	

\*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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