**Using the FRPath.org Information Input Form**

**FRPath.org**

**Where the Roads to Accelerated Assessments Converge**

1. Complete the form as best as possible. If this is being used to update an existing entry, only complete the section(s) that should be considered for updating.
2. Some responses use drop-down menus. These are indicated in red with the words “Chose an Item”. Please do not over-write these entries. If you wish to consider a response that is not available in the entry, please describe this in the final section.
3. **Please return this form to FRPath.org by email to info@frpath.org**

| *FRPath.org Country and FRP Information Input Form* | | | | | |
| --- | --- | --- | --- | --- | --- |
| Country: Click here to enter text. | | | | **Agency Name:** Click here to enter text. | |
| 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 during development | **2. Accelerates the regulatory review process** | | | | **3. Relies on or recognizes a prior regulatory decision** |
|  |  | | | |  |
| Is a Guidance or SOP describing how to apply this FRP publicly available? | | | Choose an item. | | |
| When should the FRP be requested? | | | Choose an item. | | |
| Does the agency provide assistance/advice to the sponsor? | | | Choose an item. | | |
| For which types of product(s) can this FRP be used? E.g. NMEs, generics, biologics, biosimilars, all products | | | Click here to enter text. | | |
| Must the product address an unmet medical need or serious condition? | | | Choose an item. | | |
| 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?** |
|  | |  | | |  |
| If this is a reliance or recognition pathway, what are the accepted reference agencies? | | | Click here to enter text. | | |
| How many reference agency decisions are required? | | | Click here to enter text. | | |
| Does this FRP require submission of Assessment Reports from prior decisions? | | | Choose an item. | | |
| Is a CPP (Certificate of Pharmaceutical Product) required for approval? | | | Choose an item. | | |
| Can an alternate form of reference documentation to the CPP be used? If so, what types of documents? | | | Click here to enter text. | | |
| If this process is through a Regional Regulatory Initiative, which countries participate in this process? | | | Click here to enter text. | | |
| Does the product have to have been marketed in another country? For a specific amount of time? If so, for how long? | | | Click here to enter text. | | |
| How are queries to the companies sent? | | | Choose an item. | | |
| Are external reviewers (e.g. non-agency) involved in the assessment? | | | Choose an item. | | |
| Post-authorization study commitments | | | Choose an item. | | |
| For how long is the initial approval or designation valid? | | | Choose an item. | | |
| Any other details you wish to provide? | | | Click here to enter text. | | |
| Date of this update | | |  | | |
| References | | |  | | |

\*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.

*By submitting this form, I agree that the information is true to the best of my knowledge and I consent that it can be used without restriction by FRPath.*

|  |  |
| --- | --- |
| Full Name: | Date: |
| Name of Your Affiliation: | Your Email: |
| City and Country: |  |

**Please return this form to FRPath.org by email to info@frpath.org**

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