

USP Revision Request Form

Please email the completed form to <u>donations@usp.org</u>. The donations team will review the request and create an entry within the USP Donor Portal (https://uspharm.force.com/dc/s/) for the secure sharing of support documentation.

Company Name:		
Submitter Name:		
Phone Number:		

Details for Revision Request

https://www.usp.org/get-involved/donate/submission-guidelines

Monograph Name:

Email Address:

NDA/ANDA/DMF number

(For DMF holders, please indicate the (A)NDA the DMF is referenced in)

Application Status:

For pending applications, please provide copies of any documentation from FDA recommending that you pursue the USP Pending Monograph process (you may redact sensitive information)

Comments regarding regulatory status:

GUDFA goal date (if applicable):

or Revision to the USP Requested test(s) for revision:

Is this request related to a compliance issue?:

Please provide a detailed explanation of the requested changes, to include any justification/rationale for your request. A cover letter is also acceptable: (Please also include any timing issues that USP should be aware of)

P MGs FAQ.pdf

s_Supporting Info for Disso_Drug Release_Disintegration Tests.pdf

Links to Submission Checklists and Guidelines:

Submission Checklist (the last two pages include a submission checklist)

Supporting Information for Adding Dissolution Tests

Pending Monograph Guidelines

Please note that the next step for your revision request is a series of reviews. If there are any additional data needs or questions, a USP staff member will reach out for the information required. Due to operational priorities, there may be a delay in addressing your request. In addition, after the request has been reviewed, the Documentary Standard Scientist and Expert Committee for this monograph will decide whether to pursue the revision request.