

- (i) Summaries of significant safety findings from clinical trials during the reporting period;
 - (ii) Findings from non-interventional Studies;
 - (iii) Findings from non-Clinical Studies;
 - (iv) Findings from literature.
- (i) Other information: This section of periodic safety update reports should include the details about signals and Risk Management Plan in place by licence holder (if any).
- (a) Signal and risk evaluation: In this section licence holder will provide the details of signal and risk identified during the reporting period and evaluation of signals identified during the reporting period.
 - (b) Risk management plan: In this section licence holder will provide the brief details of safety concern and necessary action taken by him to mitigate these safety concerns.
- (j) Overall Safety Evaluation: This section of periodic safety update reports should capture the overall safety evaluation of the drug based upon its risk benefit evaluation for approved indication.
- (i) Summary of safety concerns
 - (ii) Benefit evaluation
 - (iii) Benefit risk analysis evaluation
- (k) Conclusion: This section of periodic safety update reports should provide the details on the safety profile of drug and necessary action taken by the licence holder in this regards.
- (l) Appendix: The appendix includes the copy of marketing authorisation in India, copy of prescribing information, line listings with narrative of Individual Case Safety Reports (ICSR).

SIXTH SCHEDULE

(See rules 21, 22, 33, 34, 45, 47, 52, 53, 60, 67, 68, 75, 76, 80, 81, 86, 91, 97 and 98)

FEE PAYABLE FOR LICENCE, PERMISSION AND REGISTRATION CERTIFICATE

| Serial Number | Rule | Subject | In rupees Indian National Rupee (INR) except where specified in dollars (\$) |
|---------------|------|---|--|
| 01 | 21 | Application for permission to conduct clinical trial | |
| | | (i) Phase I | 3,00,000 |
| | | (ii) Phase II | 2,00,000 |
| | | (iii)Phase III | 2,00,000 |
| | | (iv) Phase IV | 2,00,000 |
| 02 | 22 | Reconsideration of application for permission to conduct clinical trial | 50,000 |
| 03 | 33 | Application for permission to conduct bioavailability or bioequivalence study | 2,00,000 |
| 04 | 34 | Reconsideration of application of permission to conduct bioavailability or bioequivalence study | 50,000 |

| | | | |
|----|----|--|------------------|
| 05 | 45 | Application for registration of bioavailability and bioequivalence study centre | 5,00,000 |
| 07 | 47 | Reconsideration of application for Registration of bioavailability and bio-equivalence study centre | 1,00,000 |
| 08 | 52 | Application for permission to manufacture new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study | 5000 per product |
| 09 | 53 | Reconsideration of application to manufacture new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study | 2000 per product |
| 10 | 59 | Application for permission to manufacture unapproved active pharmaceutical ingredient for development of formulation for test or analysis or clinical trial or bioavailability or bioequivalence study | 5000 per product |

| | | | |
|----|----|---|------|
| 11 | 60 | Reconsideration of permission to Manufacture unapproved active pharmaceutical ingredient for development of formulation for test or analysis or clinical trial or bioavailability or bioequivalence study | 2000 |
|----|----|---|------|

| | | | |
|----|----|---|------------------|
| 12 | 67 | Application for import of new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study or for examination, test and analysis | 5000 per product |
| 13 | 68 | Reconsideration of application for Import of new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study or for examination, test and analysis | 1000 |
| 14 | 75 | Application for permission to import new drug (Finished Formulation) for marketing | 5,00,000 |
| 15 | | Application for permission to import new Drug (Finished Formulation) already approved in the country for marketing | 2,00,000 |
| 16 | | Application for permission to import new drug (Active Pharmaceutical Ingredient) for marketing | 5,00,000 |

| | | | |
|----|----|--|----------|
| 17 | | Application for permission to import new drug (Active Pharmaceutical Ingredient) already approved in the country for marketing | 2,00,000 |
| 18 | | Application for permission to import approved new drug for new claims, new indication or new dosage form or new route of administration or new strength for marketing | 3,00,000 |
| 19 | | Application for permission to import fixed dose combination having one or more of the ingredients as unapproved new molecules for marketing | 5,00,000 |
| 20 | | Application for permission to import fixed Dose combination having approved ingredients for marketing | 4,00,000 |
| 21 | | Application for permission to import fixed dose combination already approved for marketing | 2,00,000 |
| 22 | | Application for permission to import fixed dose combination for new claims, new indication or new dosage form or new route of administration or new strength for marketing | 3,00,000 |
| 23 | 76 | Reconsideration of application for permission to import new drug for marketing | 50,000 |

| | | | |
|----|----|--|----------|
| 24 | | Application for permission to manufacture new drug (Finished Formulation or Active Pharmaceutical Ingredient) for sale or distribution | 5,00,000 |
| 25 | | Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) already approved in the country for sale or distribution | 2,00,000 |
| 26 | 80 | Application for permission to manufacture new drug (Finished Formulation) for sale or distribution | 5,00,000 |
| 27 | | Application for permission to manufacture new drug (Finished Formulation) already approved in the country for sale or distribution | 2,00,000 |
| 28 | | Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) for sale or distribution | 5,00,000 |

| | | | |
|----|----|--|----------|
| 29 | | Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) already approved in the country for sale or distribution | 2,00,000 |
| 30 | | Application for permission to manufacture approved new drug for new claims, new indication or new dosage form or new route of administration or new strength for sale or distribution | 3,00,000 |
| 31 | | Application for permission to manufacture fixed dose combination having one or more of the ingredients as unapproved new molecules for sale or distribution | 5,00,000 |
| 32 | 80 | Application for permission to manufacture fixed dose combination having approved ingredients for sale or distribution | 3,00,000 |
| 33 | | Application for permission to manufacture fixed dose combination already approved for sale or distribution | 2,00,000 |
| 34 | | Application for permission to manufacture fixed dose combination for new claims, new indication or new dosage form or new route of administration or new strength for sale or distribution | 3,00,000 |

| | | | |
|----|----|--|----------|
| 35 | 80 | Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) or to manufacture finished formulation | 5,00,000 |
| 36 | | Application for permission to import or to manufacture phyto-pharmaceutical drugs | 2,00,000 |
| | | Reconsideration of application for | |
| 37 | 81 | permission to manufacture new drug for sale or distribution | 50,000 |
| | | Application for Import of unapproved new | |
| 38 | 86 | drug by Government hospital and medical institution | 10,000 |
| | | Application for permission to manufacture unapproved new drug but under clinical | |
| 39 | 91 | trial, for treatment of patient of life threatening disease | 5,000 |
| 40 | 98 | Pre-submission meeting | 5,00,000 |
| 41 | 99 | Post-submission meeting | 50000 |
| 42 | - | Any other application which is not specified above | 50000 |

Note 1: No fee shall be chargeable in respect of application for conduct of clinical trial for orphan drugs as defined in clause (x) of rule 2.

Note 2: In case of application received from Micro Small Medium Enterprises (MSME) firms for conduct of clinical trial, approval of new drug and pre and post submission meeting, the fee payable shall be half of the fee specified above.