**Application for Reimbursement for Licensed Medicines – New Pharmaceutical Form or Strength**

* **Parts I, II and III:** Marketing authorization holder/applicant should fill in each fields on the form. Application is not considered valid unless all the fields have been filled in.
* **Part IV:** Filled in by The National University Hospital.

**Part I. – Basic Information**

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| 1.Please fill in the following basic information: | |
| Marketing authorization holder |  |
| Local representative |  |
| Contact name |  |
| Address |  |
| Phone number |  |
| Email address |  |

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| 2. Trade name and active ingredient. |
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| 3. What is the new pharmaceutical form? |
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| 4. What is the new strength? |
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| 5. Insert link to SmPC. |

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| 6. ATC code. |

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| 7. Indication applied for? |
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| 8. Which indications have been approved for reimbursement? |

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| 9. Dosage and pharmaceutical form? |

**Part II. – Clinical Evaluation.**

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| 1. Does new strength or new pharmaceutical form effect treatment andi if so how? |

**Part III. – Economical Evaluation.**

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| 1. Please estimate the cost of treatment per individual for the indication being applied for. What is the estimated annual cost per patient treated with the drug, based on the indication applied for. What are the premises for that conclusion, what is the dosage size, duration of treatment, etc. Refer to the information on which the plan is based. |

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| 2. Please estimate the total cost for the indication being applied for. What is the estimated total annual cost for the indication being applied for the next three years, i.e. what is the number of patients, when in the year does treatment start, when does it end, what is the dosage size, etc. Refer to he information on which the plan is based. |

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| 3. Estimated cost effects on other treatments:  a. Which drugs are in use today for the same indication and in which dosage sizes. Please cite sources of the information.  b. Is it estimated that treatment and cost of other drugs with the same indication will be affected with the approval of reimbursementand and if so how? |

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| 4. Is it estimated that approval of the drug will have direct effect on other interventions/treatments within the health care service? |

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| 5. Are other benefits or changes expected should this indication be approved, such as regarding out-patients clinics, testing, blood sampling etc. |

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| 6. Overview of submitted health economical evaluation, CEA (Cost Effective Analysis) | | | |
| Name/Country | Type | Indication | Comparator |
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| 7. What effect do the results of the CEA have on the cost of treatment with the drug, and especially those that have been conducted in the reference countries? | | | |

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| 8.a. Is the indication in the CEA consistent with the estimated use in Iceland?  8.b. Are the patient groups the same in the CEA? |

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| 9. List an overview of reimbursements in reference countries. Please state whether it is general, with conditions and if so which, or individual. Please insert links to information from reference countries. | | | | |
| Reference country | General reimbursement | Conditions | Individual reimbursement | Link |
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| 10. Other aspects of economical evaluation for consideration? |

**Part IV. – Assessment by Icelandic Medicine Pricing and Reimbursement Committee at The National University Hospital of Iceland (Lyfjanefnd Landspítala) (IMPRC)**

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| 1. IMPRC‘s full assessment: |

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| 2. Summary of clinical and economical evaluation by IMPRC: |

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| 3. Date of summary/by: |