| CHI no | | | |
|------------|------|---|-----|
| First name | DOB | | |
| Last name | Sex: | M | 🗌 F |
| Address | | | |
| | | | |

Service/Hospitals/Dept. etc. Ward/Team:



Appendix 2 - Blanket Unlicensed & High Risk Off Label Medicine Application

or attach addressograph label here

Date: Time:

Time: (24 hour)

Application Form

Identifies as _____

This form is to be used in conjunction with the NHS Lanarkshire Policy for Unlicensed Medicines. Before completion, you must have read this policy which identifies your responsibilities.

| Requester detai | ls | | | | |
|---|---|--|---|--|--|
| Prescriber name: | | Hospital site: | | | |
| Speciality: | | Ward/Outpatient dept: | | | |
| Contact details: | | Date requested: | | | |
| | | Date required: | | | |
| Patient details | | | | | |
| Anticipated usag | e (please tick) - Estimated patient numb | pers: | | | |
| For your patie | ents only |] For patients within your speciality on a single site | | | |
| For patients w | vithin your speciality on all sites |] Any patient within the Health Board | | | |
| Unlicensed Mec | licine Details | | | | |
| Product name: (International No | on Proprietary Name) | | | | |
| Proprietary Nam | | | | | |
| Strength and Pha | armaceutical Form: | | | | |
| Manufacturer (if | <nown):< td=""><td></td><td></td></nown):<> | | | | |
| Indication: | | | | | |
| Dose/frequency/ | route: | | | | |
| Duration of Treat | ment: | | | | |
| Category of req | uest: | | | | |
| | | narketing authorisation for a licensed medicine | _ | | |
| (off-label prescribing) and is considered 'high risk' in Appendix 4 | | | | | |
| | | | | | |
| | is unlicensed – please complete the fo nsed medicine being considered? (Tick | - | | | |
| - | IK licensed product available to treat or | | | | |
| 2. The UK licensed product used to treat or diagnose the medical condition is temporarily unavailable | | | | | |
| | sed product used to treat or diagnose t | | | | |
| 4. No therapeutically equivalent UK licensed product available or suitable (provide details): | | | | | |
| 5. Patient Safety: | | | | | |
| 6. Other (provid | le details): | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| 5 (Apr | | | | | |
| ot | | | | | |
| Was a product licence in the UK withdrawn? 🗌 Yes 📄 No 📄 Not known | | | | | |

NMAHP.BU&HTO.24_24515.L

Blanket Unlicensed & High Risk Off Label Medicine | NHSL | Page 1 of 3

If yes, contact manufacturer to find out reasons for withdrawal.

| Patient name: | CHI number: | | | |
|--|-------------------------|------------|----------|-------------|
| Clinical Evidence | | | | |
| Is there any evidence to support its use for the proposed | indication? | 🗌 Yes | 🗌 No | |
| Is there evidence to support its proposed administration (dose, duration, concentration for parenteral products an | | 🗌 Yes | 🗌 No | |
| Is the active drug currently in a licensed product for use v of administration e.g. tablet, suspension? | ia the same route | 🗌 Yes | 🗌 No | |
| Is the product licensed for the specified indication in ano | ther country? | 🗌 Yes | 🗌 No | 🗌 Not known |
| UK product licence applied for? If yes, record date of application for licence: | | 🗌 Yes | 🗌 No | 🗌 Not known |
| Are other Boards using this medicine? If so, name: | | 🗌 Yes | 🗌 No | 🗌 Not known |
| Summarise below the supporting evidence, list reference | s and attach copies of | reference | es where | available. |
| | | | | |
| What side effects and significant interactions have been r | | | | |
| Give details of contraindications and any other risks to the | e patient. Include pred | cautions i | n use. | |
| | - Panerer merere hre | | | |
| Will there be any primary care implications? (e.g. need for a shared care protocol) If so, describe: | | | | |
| | | | | |

| Patient name: CHI nun | nber: |
|-----------------------|-------|
| | |

| Prescriber | (SpR) GP or | other prescr | iber (| Tick one) | |
|--|-------------------|---------------------|-------------------------|----------------------|--|
| Print name: | | | Speciality/Directorate: | | |
| Signature: | | Date: | | | |
| If SpR, state name of patient's consultar | nt: | 1 | | | |
| Authorisation of Application (pharma | cy – acute senior | pharmacist or lo | ocality | prescribing adviser) | |
| Name | Designation | | Signature & Date | | |
| | | | | | |
| Medicines Cost (Medicines costing less processes in primary care or go straight | | | | | |
| For medicines costing more than £5,00 but less than £25,000 per patient/year? Approved by acute site Chief of Medici | Yes 🗌 No | Director (Primary (| Care) | Signature | |
| For medicines costing more than £25,0 | 00 | | Signature | | |
| per patient/year? Yes No Approved by acute site Chief of Medicine AND Medical Director, or Associate Director (Primary Care) | | Director, | Signature | | |
| Final process approval | | | | | |
| Approval for use 🗌 Yes 🗌 No | Date: | | | | |
| If no, give reasons | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| State restrictions on prescribing/use | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| Any further Information | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |
| Completed by: (PRINT NAME) | D | esignation of ap | prove | r: | |
| Signature: | | | | | |
| | D | ate: | | Time: | |