Use this checklist to determine if a patient meets the restrictions for funding in the **hospital setting**. For more details, refer to Section H of the Pharmaceutical Schedule. For community funding, see the Special Authority Criteria.

| PRESCR         | IBER                    | PATIENT:   |
|----------------|-------------------------|--|
| Name:          |                         |  |
| Ward:          |                         | NHI:   |
| Laronidase     |                         |  |
|                | ssmer<br>isites<br>Pres | nt required after 24 weeks (tick boxes where appropriate) cribed by, or recommended by a metabolic physician, or in accordance with a protocol or guideline that has been endorsed by the Health dospital.   |
| and            | O                       | The patient has been diagnosed with Hurler Syndrome (mucopolysacchardosis I-H)   |
|                | or                      | O Diagnosis confirmed by demonstration of alpha-L-iduronidase deficiency in white blood cells by either enzyme assay in cultured skin fibroblasts O Detection of two disease causing mutations in the alpha-L-iduronidase gene and patient has a sibling who is known to have Hurler syndrome                      |
| an<br>an<br>an | d<br>O                  | Patient is going to proceed with a haematopoietic stem cell transplant (HSCT) within the next 3 months and treatment with laronidase would be bridging treatment to transplant  Patient has not required long-term invasive ventilation for respiratory failure prior to starting Enzyme Replacement Therapy (ERT) |
|                | <u> </u>                | Laronidase to be administered for a total of 24 weeks (equivalent to 12 weeks pre- and 12 post-HSCT) at doses no greater than 100 units/kg every week  |