Enquiries to Ministry of Health 0800 855 066

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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| PLICANT (stamp or sticker acceptable) | | | PATIENT NHI: | REFERRER Reg No: |
|---------------------------------------|--|--|---|---|
| ۷o: | | | First Names: | First Names: |
| me: | | | Surname: | Surname: |
| ress: | | | DOB: | Address: |
| | | | Address: | |
| lumber | ······································ | | | Fax Number: |
| ximat | o (Ma | othera) | | |
| lications | s only | n — rheumatoid arthritis - TNF from a rheumatologist or Practit k boxes where appropriate) | Finhibitors contraindicated ioner on the recommendation of a rheumatologist. Ap | provals valid for 4 months. |
| and | T | reatment with a Tumour Necrosis | s Factor alpha inhibitor is contraindicated | |
| | | | e erosive rheumatoid arthritis (either confirmed by radi | ology imaging, or the patient is cyclic citrullinated |
| and | 一 ['] | , , , , , , | for six months duration or longer | |
| L | | atient has tried and not respond naximum tolerated dose | ed to at least three months of oral or parenteral metho | trexate at a dose of at least 20 mg weekly or a |
| and | | | ed to at least three months of oral or parenteral metho | trexate in combination with sulfasalazine and |
| and | h | ydroxychloroquine sulphate (at n | naximum tolerated doses) | |
| | or [| Patient has tried and not re tolerated dose of ciclosporia | sponded to at least three months of oral or parenteral n | methotrexate in combination with the maximum |
| | or [| Patient has tried and not regold | sponded to at least three months of oral or parenteral | methotrexate in combination with intramuscular |
| |) | Patient has tried and not re combination with oral or pa | sponded to at least three months of therapy at the ma renteral methotrexate | ximum tolerated dose of leflunomide alone or in |
| and | | _ | | |
| | or | Patient has persistent symp | otoms of poorly controlled and active disease in at least | st 20 swollen, tender joints |
| | | Patient has persistent symp knee, ankle, and either sho | otoms of poorly controlled and active disease in at leasulder or hip | st four joints from the following: wrist, elbow, |
| and | ı | | | |
| | or | Patient has a C-reactive pro | otein level greater than 15 mg/L measured no more that | an one month prior to the date of this application |
| | | C-reactive protein levels no day and has done so for mo | t measured as patient is currently receiving prednison ore than three months | e therapy at a dose of greater than 5 mg per |
| and | [| Rituximab to be used as an | adjunct to methotrexate or leflunomide therapy | |
| | or | | b both methotrexate and leflunomide, requiring rituxima | ah monotherany to be used |
| | I | | , pour mounding and and ionarionnes, requiring maximi | ab interioritierapy to be asea |

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| | Address: | | | | | | | | |
| | | | | | | | | | |
| Fax Number: | | Fax Number: | | | | | | | |
| Rituximab (Mabthera) - continued | | | | | | | | | |
| Initial application — rheumatoid arthritis - prior Applications only from a rheumatologist or Practitio Prerequisites(tick boxes where appropriate) | TNF inhibitor use ner on the recommendation of a rheumatologist. App | provals valid for 4 months. | | | | | | | |
| The patient has had an initial rheumatoid arthritis | | | | | | | | | |
| | enced intolerable side effects from a reasonable trial | of adalimumab and/or etanercept | | | | | | | |
| Following at least a fou | Following at least a four month trial of adalimumab and/or etanercept, the patient did not meet the renewal criteria for adalimumab and/or etanercept for rheumatoid arthritis | | | | | | | | |
| and | | | | | | | | | |
| | Rituximab to be used as an adjunct to methotrexate or leflunomide therapy | | | | | | | | |
| or Patient is contraindicated to be | both methotrexate and leflunomide, requiring rituxima | ab monotherapy to be used | | | | | | | |
| and Maximum of two 1,000 mg infusion | | | | | | | | | |
| | | | | | | | | | |
| Renewal — rheumatoid arthritis - re-treatment in | | | | | | | | | |
| Current approval Number (if known): Applications only from a rheumatologist or Practitio | ner on the recommendation of a rheumatologist. Ap | provals valid for 4 months. | | | | | | | |
| Prerequisites(tick boxes where appropriate) | , | | | | | | | | |
| | tial course of rituximab infusions the patient had betwinically significant response to treatment in the opinio | | | | | | | | |
| | cond course of rituximab infusions the patient had at ifficant response to treatment in the opinion of the phy | | | | | | | | |
| At 4 months following the thin | rd and subsequent courses of rituximab infusions, the point count from baseline and a clinically significant res | | | | | | | | |
| and Rituximab re-treatment not to be gir | ven within 6 months of the previous course of treatme | ent | | | | | | | |
| | adjunct to methotrexate or leflunomide therapy | | | | | | | | |
| or Patient is contraindicated to be | both methotrexate and leflunomide, requiring rituxima | ub monotherapy to be used | | | | | | | |
| and Maximum of two 1,000 mg infusion | s of rituximab given two weeks apart | | | | | | | | |

I confirm the above details are correct and that in signing this form I understand I may be audited.

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| | Address: | | | | | | | | |
| | | | | | | | | | |
| Fax Number: | | Fax Number: | | | | | | | |
| Rituximab (Mabthera) - continued | | | | | | | | | |
| Applications only from a rheumatologist or Practition Prerequisites (tick boxes where appropriate) At 4 months following the inition baseline and a clinically sign or At 4 months following the se 30% improvement in active juphysician and Rituximab re-treatment not to be grand Rituximab to be used as an active juphysician | At 4 months following the initial course of rituximab infusions the patient had at least a 50% decrease in active joint count from baseline and a clinically significant response to treatment in the opinion of the physician At 4 months following the second and subsequent courses of rituximab infusions, the patient demonstrates at least a continuing 30% improvement in active joint count from baseline and a clinically significant response to treatment in the opinion of the | | | | | | | | |
| and Maximum of two 1,000 mg infusion | ns of rituximab given two weeks apart | | | | | | | | |

I confirm the above details are correct and that in signing this form I understand I may be audited.