RS2133 - Rituximab

		1
ABO-incompatible organ transplant - INITIATION	9	l
ANCA associated vasculitis - INITIATION	8	l
ANCA associated vasculitis - CONTINUATION	8	l
Antibody-mediated organ transplant rejection - INITIATION		l
B-cell acute lymphoblastic leukaemia/lymphoma* - INITIATION	.16	l
CD20+ low grade or follicular B-cell NHL - INITIATION		ı
CD20+ low grade or follicular B-cell NHL - CONTINUATION		l
Chronic lymphocytic leukaemia - INITIATION	4	l
Chronic lymphocytic leukaemia - CONTINUATION	5	l
Membranous nephropathy - INITIATION	.15	l
Membranous nephropathy - CONTINUATION	.15	l
Neuromyelitis Optica Spectrum Disorder (NMOSD) - INITIATION	.11	l
Neuromyelitis Optica Spectrum Disorder (NMOSD) - CONTINUATION	.11	l
Severe Refractory Myasthenia Gravis - INITIATION	.12	l
Severe Refractory Myasthenia Gravis - CONTINUATION	.12	l
Severe antisynthetase syndrome - INITIATION	.12	l
Severe antisynthetase syndrome - CONTINUATION	.13	l
Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) - INITIATI	ON	l
10		l
Steroid dependent nephrotic syndrome (SDNS) or frequently relapsing nephrotic syndrome (FRNS) - CONTINU.	ATIO	۱
10		l
Steroid resistant nephrotic syndrome (SRNS) - INITIATION	.10	l
Steroid resistant nephrotic syndrome (SRNS) - CONTINUATION	.11	l
Aggressive CD20 positive NHL - INITIATION		l
Aggressive CD20 positive NHL - CONTINUATION	4	l
Anti-NMDA receptor autoimmune encephalitis - INITIATION	.14	l
Anti-NMDA receptor autoimmune encephalitis - CONTINUATION	.14	l
Desensitisation prior to transplant - INITIATION	.16	l
Graft versus host disease - INITIATION		l
Haemophilia with inhibitors - INITIATION	2	l
Haemophilia with inhibitors - CONTINUATION	2	l
Immune thrombocytopenic purpura (ITP) - INITIATION	6	l
Immune thrombocytopenic purpura (ITP) - CONTINUATION	7	l
Immunoglobulin G4-related disease (IgG4-RD*) - INITIATION	.17	l
Immunoglobulin G4-related disease (IgG4-RD*) - CONTINUATION	.17	١
Indolent, low-grade lymphomas or hairy cell leukaemia* - INITIATION	3	١
Indolent, low-grade lymphomas or hairy cell leukaemia* - CONTINUATION	3	l
Pemiphigus* - INITIATION		l
Pemiphigus* - CONTINUATION		l
Post-transplant - INITIATION	2	l
Post-transplant - CONTINUATION	2	l
Pure red cell aplasia (PRCA) - INITIATION	8	l
Pure red cell aplasia (PRCA) - CONTINUATION		l
Severe chronic inflammatory demyelinating polyneuropathy - INITIATION	.13	l
Severe chronic inflammatory demyelinating polyneuropathy - CONTINUATION	.13	
Severe cold haemagglutinin disease (CHAD) - INITIATION	5	
Severe cold haemagglutinin disease (CHAD) - CONTINUATION	5	
Thrombotic thrombocytopenic purpura (TTP) - INITIATION		1
Thrombotic thrombocytopenic purpura (TTP) - CONTINUATION		1
Treatment refractory systemic lupus erythematosus (SLE) - INITIATION	9	
Treatment refractory systemic lupus erythematosus (SLE) - CONTINUATION	9	۱
Warm autoimmune haemolytic anaemia (warm AIHA) - INITIATION	6	1
Warm autoimmune haemolytic anaemia (warm AIHA) - CONTINUATION	6	1

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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.

PRESCRIBER PATIENT:		PATIENT:	
Name: Name:		Name:	
Ward:		NHI:	
Rituxin	nab (Riximyo)		
	ION – haemophilia with inhibitors uisites (tick boxes where appropriate)		
and	Prescribed by, or recommended by a haematologist, or in accordance Hospital.	e with a protocol or guideline that has been endorsed by the Health NZ	
OI	O Patient has mild congenital haemophilia complicated by inhibite	prs	
OI	O Patient has severe congenital haemophilia complicated by inhil	bitors and has failed immune tolerance therapy	
	O Patient has acquired haemophilia		
		e with a protocol or guideline that has been endorsed by the Health NZ	
and ar	O An initial response lasting at least 12 months was demonstrate		
	ION – post-transplant uisites (tick boxes where appropriate)		
ar	The patient has B-cell post-transplant lymphoproliferative disorned To be used for a maximum of 8 treatment cycles	der*	
Note: Ir	dications marked with * are unapproved indications.		
	CONTINUATION – post-transplant Prerequisites (tick boxes where appropriate)		
ar	O The patient has B-cell post-transplant lymphoproliferative disor		
Note: Ir	dications marked with * are unapproved indications.		

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.

PRESCRIBER			PATIENT:	
Name:			Name:	
Ward:				NHI:
Ritu	kima	ı b (Rixin	nyo) - continued	
Re-a	ssess	ment re	elent, low-grade lymphomas or hairy cell leukaemia* quired after 9 months k boxes where appropriate)	
	or	and	The patient has indolent low grade NHL or hairy cell leu To be used for a maximum of 6 treatment cycles	kaemia* with relapsed disease following prior chemotherapy
	OI.	and	The patient has indolent, low grade lymphoma or hairy To be used for a maximum of 6 treatment cycles	cell leukaemia* requiring first-line systemic chemotherapy
			v-grade lymphomas' includes follicular, mantle, marginal zo ell leukaemia' also includes hairy cell leukaemia variant.	one and lymphoplasmacytic/Waldenstrom macroglobulinaemia. *Unapproved
Re-a	CONTINUATION – indolent, low-grade lymphomas or hairy cell leukaemia* Re-assessment required after 12 months Prerequisites (tick boxes where appropriate) The patient has had a rituximab treatment-free interval of 12 months or more and The patient has indolent, low-grade NHL or hairy cell leukaemia* with relapsed disease following prior chemotherapy and To be used for no more than 6 treatment cycles Note: 'Indolent, low-grade lymphomas' includes follicular, mantle, marginal zone and lymphoplasmacytic/Waldenstrom macroglobulinaemia. *Unapproved indication. 'Hairy cell leukaemia' also includes hairy cell leukaemia variant.			
			ressive CD20 positive NHL s boxes where appropriate)	
	or	and and and	The patient has treatment naive aggressive CD20 position To be used with a multi-agent chemotherapy regimen given the patient for a maximum of 8 treatment cycles The patient has aggressive CD20 positive NHL with relation to be used for a maximum of 6 treatment cycles	even with curative intent
Note	: 'Aan	ressive	CD20 positive NHL' includes large B-cell lymphoma and Bu	urkitt's lymphoma/leukaemia.
11018	. , 199	,. CGGIVG	DEED POSITION IN IL MONIGODO RAIGO D COM TYMPHOMA AND DE	

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.

PRESCRIBER		PATIENT:	
Name:		Name:	
Ward:		NHI:	
Rituximab (Riximyo) - continued		
	ON – aggressive CD20 positive NHL s (tick boxes where appropriate)		
and and and and	The patient has had a rituximab treatment-free interval of 12 n. The patient has relapsed refractory/aggressive CD20 positive. To be used with a multi-agent chemotherapy regimen given with the compact of the compac	NHL ith curative intent	
		initio iyinpioina earaeinia.	
Re-assessmer	Chronic lymphocytic leukaemia ent required after 12 months s (tick boxes where appropriate)		
and	The patient has progressive Binet stage A, B or C chronic lym	phocytic leukaemia (CLL) requiring treatment	
or	The patient is rituximab treatment naive		
	or The patient is chemotherapy treatment naive		
	and	g no more than three prior lines of chemotherapy treatment al of 12 months or more if previously treated with fludarabine and	
or		nent is to be used in combination with funded venetoclax	
The patient has good performance status and			
	O The patient does not have chromosome 17p deletion CL		
or		unded venetoclax for relapsed/refractory chronic lymphocytic leukaemia	
and	Rituximab to be administered in combination with fludarabine 6 treatment cycles	and cyclophosphamide, bendamustine or venetoclax for a maximum of	
	It is planned that the patient receives full dose fludarabine and bendamustine or venetoclax	d cyclophosphamide (orally or dose equivalent intravenous administration),	
standard thera temporarily de	apeutic chemotherapy regimen and supportive treatments. 'Goo	shoma. A line of chemotherapy treatment is considered to comprise a known of performance status' means ECOG score of 0-1, however, in patients 3) is acceptable where treatment with rituximab is expected to improve	

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

PRESCRIBER	PATIENT:		
ame: Name:			
/ard:NHI:			
Rituximab (Riximyo) - continued			
CONTINUATION – Chronic lymphocytic leukaemia Re-assessment required after 12 months			
Prerequisites (tick boxes where appropriate)			
O The patient's disease has relapsed and ritux	timab treatment is to be used in combination with funded venetoclax		
The patient's disease has relapsed foll	lowing no more than one prior line of treatment with rituximab for CLL		
	months or more since commencement of initial rituximab treatment		
The patient does not have chromosom	ne 17p deletion CLL		
It is planned that the patient receives for administration) or bendamustin	full dose fludarabine and cyclophosphamide (orally or dose equivalent intravenous		
and Rituximab to be administered in combination with f 6 treatment cycles	fludarabine and cyclophosphamide, bendamustine or venetoclax for a maximum of		
·	nocytic lymphoma. A line of chemotherapy treatment is considered to comprise a known nents.		
And O Patient has cold haemagglutinin disease* and O Patient has severe disease which is characterized symptoms	n accordance with a protocol or guideline that has been endorsed by the Health NZ by symptomatic anaemia, transfusion dependence or disabling circulatory		
The total rituximab dose used would not exceed th	e equivalent of 375 mg/m2 of body surface area per week for a total of 4 weeks		
Note: Indications marked with * are unapproved indications.			
CONTINUATION – severe cold haemagglutinin disease (CHAD Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate)))		
O Prescribed by, or recommended by a haematologist, or in Hospital.	n accordance with a protocol or guideline that has been endorsed by the Health NZ		
O Previous treatment with lower doses of rituximab (doses (375 mg/m² weekly for 4 weeks) is now plan or	100 mg weekly for 4 weeks) have proven ineffective and treatment with higher nned		
Patient was previously treated with rituximab	o for severe cold haemagglutinin disease*		
An initial response lasting at least 12 months and	s was demonstrated		
O Patient now requires repeat treatment			
Note: Indications marked with * are unapproved indications.			

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PRESCRIBER	PATIENT:			
Name:	Name:			
Ward:NHI:				
Rituximab (Riximyo) - continued				
INITIATION – warm autoimmune haemolytic anaemia (warm AIHA) Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate)				
Prescribed by, or recommended by a haematologist, or in accordance Hospital.	e with a protocol or guideline that has been endorsed by the Health NZ			
> 5 mg prednisone daily), cytotoxic agents (e.g. cyclophospha	(including if patient requires ongoing steroids at doses equivalent to mide monotherapy or in combination), intravenous immunoglobulin tof 375 mg/m2 of body surface area per week for a total of 4 weeks			
Note: Indications marked with * are unapproved indications.				
CONTINUATION – warm autoimmune haemolytic anaemia (warm AIHA) Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate)				
Prescribed by, or recommended by a haematologist, or in accordance Hospital.	e with a protocol or guideline that has been endorsed by the Health NZ			
	ekly for 4 weeks) have proven ineffective and treatment with higher			
O Patient was previously treated with rituximab for warm a	utoimmune haemolytic anaemia*			
An initial response lasting at least 12 months was demon	nstrated			
Patient now requires repeat treatment				
Note: Indications marked with * are unapproved indications.				
INITIATION – immune thrombocytopenic purpura (ITP) Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a haematologist, or in accordance Hospital.	e with a protocol or guideline that has been endorsed by the Health NZ			
O Patient has immune thrombocytopenic purpura* with a p	latelet count of less than or equal to 20,000 platelets per microlitre			
Patient has immune thrombocytopenic purpura* with a p mucocutaneous bleeding	latelet count of 20,000 to 30,000 platelets per microlitre and significant			
and				
Treatment with steroids and splenectomy have been ine	ffective			
O Treatment with steroids has been ineffective and splene	ctomy is an absolute contraindication			
	e and patient is being prepared for elective surgery (e.g. splenectomy)			
The total rituximab dose used would not exceed the equivalen	t of 375 mg/m2 of body surface area per week for a total of 4 weeks			
Note: Indications marked with * are unapproved indications.				
I confirm that the above details are correct:				

Signed: Date:

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PRESCRIBER		PATIENT:		
Name: Name:		Name:		
Ward: NHI:				
Rituxin	nab (Riximyo) - continued			
Re-asse	IUATION – immune thrombocytopenic purpura (ITP) ISSMENT required after 8 weeks IIISITES (tick boxes where appropriate) Prescribed by, or recommended by a haematologist, or in acco Hospital.	rdance with a protocol or guideline that has been endorsed by the Health NZ		
and	doses (375 mg/m ² weekly for 4 weeks) is now planned	g weekly for 4 weeks) have proven ineffective and treatment with higher		
	Patient was previously treated with rituximab for im and An initial response lasting at least 12 months was cand Patient now requires repeat treatment			
Note: In	dications marked with * are unapproved indications.			
and	Hospital. The total rituximab dose used would not exceed the equinal Patient has thrombotic thrombocytopenic purpura* thrombocytopenia despite plasma exchange	rdance with a protocol or guideline that has been endorsed by the Health NZ valent of 375 mg/m2 of body surface area per week for a total of 4 weeks and has experienced progression of clinical symptoms or persistent openic purpura* with neurological or cardiovascular pathology		
Note: In	dications marked with * are unapproved indications.			
Re-asse	IUATION – thrombotic thrombocytopenic purpura (TTP) issment required after 8 weeks iisites (tick boxes where appropriate) Prescribed by, or recommended by a haematologist, or in acco Hospital.	rdance with a protocol or guideline that has been endorsed by the Health NZ		
ar	O An initial response lasting at least 12 months was demon			
ar	Patient now requires repeat treatment	valent of 375 mg/m2 of body surface area per week for a total of 4 weeks		
Nata In	Note: Indications marked with * are unapproved indications.			
Note: in	idications marked with lare unapproved indications.			

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.

PRESCRIBER	PATIENT:		
Name:	Name:		
Ward:	NHI:		
Rituximab (Riximyo) - continued			
INITIATION – pure red cell aplasia (PRCA) Re-assessment required after 6 weeks Prerequisites (tick box where appropriate) O Prescribed by, or recommended by a haematologist, or in accordance Hospital. and O Patient has autoimmune pure red cell aplasia* associated with a dem Note: Indications marked with * are unapproved indications.	e with a protocol or guideline that has been endorsed by the Health NZ nonstrable B-cell lymphoproliferative disorder		
CONTINUATION – pure red cell aplasia (PRCA) Re-assessment required after 6 weeks Prerequisites (tick box where appropriate) O Prescribed by, or recommended by a haematologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital. and O Patient was previously treated with rituximab for pure red cell aplasia* associated with a demonstrable B-cell lymphoproliferative disorder and demonstrated an initial response lasting at least 12 months Note: Indications marked with * are unapproved indications.			
INITIATION – ANCA associated vasculitis Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate)			
or disease after at least 3 months	clophosphamide has failed to achieve significant improvement of osphamide > 15 g or a further repeat 3 month induction course of 5 g		
CONTINUATION – ANCA associated vasculitis Re-assessment required after 8 weeks Prerequisites (tick boxes where appropriate)			
Patient has been diagnosed with ANCA associated vasculitis* O Patient has previously responded to treatment with rituximab be and The total rituximab dose would not exceed the equivalent of 37 Note: Indications marked with * are unapproved indications.			

I confirm that the above details are correct:

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PRESCRIBER	PATIENT:	
Name:	Name:	
Ward:	NHI:	
Rituximab (Riximyo) - continued		
the Health NZ Hospital. The patient has severe, immediately life- or organ-threatening and The disease has proved refractory to treatment with steroids a and		
mofetil and high dose cyclophosphamide, or cyclophosphamid and Maximum of four 1000 mg infusions of rituximab Note: Indications marked with * are unapproved indications.		
CONTINUATION – treatment refractory systemic lupus erythematosus (SLE) Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a rheumatologist or nephrologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ Hospital. and O Patient's SLE* achieved at least a partial response to the previous round of prior rituximab treatment and O The disease has subsequently relapsed and O Maximum of two 1000 mg infusions of rituximab Note: Indications marked with * are unapproved indications.		
INITIATION – Antibody-mediated organ transplant rejection Prerequisites (tick box where appropriate) O Patient has been diagnosed with antibody-mediated organ transplant rejection* Note: Indications marked with * are unapproved indications.		
INITIATION – ABO-incompatible organ transplant Prerequisites (tick box where appropriate) O Patient is to undergo an ABO-incompatible solid organ transplant* Note: Indications marked with * are unapproved indications.		

1	confirm that the above details are correct:	
,	Signed:	Date:

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

August 2025

PRES	CRIB	ER	PATIENT	Т:
Name			Name:	
Ward:			NHI:	
Ritux	imal	b (R	(Riximyo) - continued	
			- Steroid dependent nephrotic syndrome (SDNS) or frequently relapsion trequired after 8 weeks	ing nephrotic syndrome (FRNS)
			s (tick boxes where appropriate)	
and			scribed by, or recommended by a nephrologist, or in accordance with a prospital.	otocol or guideline that has been endorsed by the Health NZ
	(C	Patient is a child with SDNS* or FRNS*	
	and (C	Treatment with steroids for at least a period of 3 months has been ineffe	ective or associated with evidence of steroid toxicity
	and (C	Treatment with ciclosporin for at least a period of 3 months has been inc	effective and/or discontinued due to unacceptable side effects
	and (C	Treatment with mycophenolate for at least a period of 3 months with no	reduction in disease relapses
	(C	The total rituximab dose used would not exceed the equivalent of 375 m	mg/m² of body surface area per week for a total of 4 weeks
Note:	Indic	atio	ions marked with a * are unapproved indications.	
and	equisi P H and and	resolospi	ent required after 8 weeks s (tick boxes where appropriate) scribed by, or recommended by a nephrologist, or in accordance with a prospital. Patient who was previously treated with rituximab for nephrotic syndrom Treatment with rituximab was previously successful and has demonstrat relapsed and the patient now requires repeat treatment The total rituximab dose used would not exceed the equivalent of 375 in	ted sustained response for > 6 months, but the condition has
Note:	Indic	atio	ions marked with a * are unapproved indications.	
Re-as	ssessr e quisi P	men tes	- Steroid resistant nephrotic syndrome (SRNS) ent required after 8 weeks s (tick boxes where appropriate) scribed by, or recommended by a nephrologist, or in accordance with a prospital.	otocol or guideline that has been endorsed by the Health NZ
	and	C	Patient is a child with SRNS* where treatment with steroids and ciclospo	orin for at least 3 months have been ineffective
	and)	Treatment with tacrolimus for at least 3 months has been ineffective	
	and	C	Genetic causes of nephrotic syndrome have been excluded	
		<u>Э</u>	The total rituximab dose used would not exceed the equivalent of 375 in	ng/m² of body surface area per week for a total of 4 weeks
Note:	Indic	atio	ions marked with a * are unapproved indications.	

August 2025

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PRESCRIBER	PATIENT:
Name:	
Nard:NHI:	
Rituximab (Riximyo) - continued	
Hospital. Patient who was previously treated with rituximab for nephrotic and Treatment with rituximab was previously successful and has decondition has relapsed and the patient now requires repeat treatment.	emonstrated sustained response for greater than 6 months, but the
Note: Indications marked with a * are unapproved indications.	
weekly for four weeks	olate
CONTINUATION – Neuromyelitis Optica Spectrum Disorder (NMOSD) Re-assessment required after 2 years Prerequisites (tick boxes where appropriate)	
One of the following dose regimens is to be used: 2 doses of weekly for four weeks and The patients has responded to the most recent course of rituxiand The patient has not received rituximab in the previous 6 month	

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:
Rituximab (Riximyo) - continued	
INITIATION – Severe Refractory Myasthenia Gravis Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) Prescribed by, or recommended by a neurologist, or in accordance will Hospital.	vith a protocol or guideline that has been endorsed by the Health NZ
and	of body surface area per week for a total of four weeks, or 500 mg once s apart
or ineffective	munosuppressant for at least a period of 12 months has been
Treatment with at least one other immunosuppress and Corticosteroids have been trialed for at least 12 m	onths and have been discontinued due to unacceptable side effects
CONTINUATION – Severe Refractory Myasthenia Gravis Re-assessment required after 2 years Prerequisites (tick boxes where appropriate) Oreof the following dose regimens is to be used: 375 mg/m2 weekly for four weeks, or two 1,000 mg doses given two week and An initial response lasting at least 12 months was demonstrate and	of body surface area per week for a total of four weeks, or 500 mg once s apart
or least 12 months The patient's myasthenia gravis has relapsed desplaced least 12 months	onths and have been discontinued due to unacceptable side effects
INITIATION – Severe antisynthetase syndrome	
Re-assessment required after 12 months Prerequisites (tick boxes where appropriate)	
Patient has confirmed antisynthetase syndrome and Patient has severe, immediately life or organ threatening disea and Treatment with at least 3 immunosuppressants (oral ste azathioprine) has not be effective at controlling active disease.	roids, cyclophosphamide, methotrexate, mycophenolate, ciclosporin,
And Maximum of four 1,000 mg infusions of rituximab	lications

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

	Name:
Ward:	
	NHI:
Rituximab (Ri	ximyo) - continued
Re-assessment	N – Severe antisynthetase syndrome required after 12 months tick boxes where appropriate)
and and	Patient's disease has responded to the previous rituximab treatment with demonstrated improvement in inflammatory markers, muscle strength and pulmonary function The patient has not received rituximab in the previous 6 months Maximum of two cycles of 2 × 1,000 mg infusions of rituximab given two weeks apart
_	raft versus host disease tick boxes where appropriate)
and -	Patient has refractory graft versus host disease following transplant Treatment with at least 3 immunosuppressants (oral steroids, ciclosporin, tacrolimus, mycophenolate, sirolimus) has not be effective at controlling active disease
and	The total rituximab dose used would not exceed the equivalent of 375 mg/m² of body surface area per week for a total of 4 weeks
Prescr Hospit	tick boxes where appropriate) ribed by, or recommended by a neurologist, or in accordance with a protocol or guideline that has been endorsed by the Health NZ ital. Patient has severe chronic inflammatory demyelinating polyneuropathy (CIPD)
and	Treatment with steroids and intravenous immunoglobulin and/or plasma exchange has not been effective at controlling active disease At least one other immunosuppressant (cyclophosphamide, ciclosporin, tacrolimus, mycophenolate) has not been effective at controlling active disease
	Rapid treatment is required due to life threatening complications One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart
Re-assessment	N – severe chronic inflammatory demyelinating polyneuropathy required after 6 months tick boxes where appropriate)
and	Patient's disease has responded to the previous rituximab treatment with demonstrated improvement in neurological function compared to baseline
and	The patient has not received rituximab in the previous 6 months One of the following dose regimens is to be used: 375 mg/m2 of body surface area per week for a total of four weeks, or 500 mg once weekly for four weeks, or two 1,000 mg doses given two weeks apart

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PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:
Rituximab (Riximyo) - continued	
INITIATION – anti-NMDA receptor autoimmune encephalitis Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) O Prescribed by, or recommended by a neurologist, or in accordance Hospital.	with a protocol or guideline that has been endorsed by the Health NZ
Patient has severe anti-NMDA receptor autoimmune encepha	itis
active disease At least one other immunosuppressant (cyclophose effective at controlling active disease	lobulin and/or plasma exchange has not been effective at controlling sphamide, ciclosporin, tacrolimus, mycophenolate) has not been
Rapid treatment is required due to life threatening compand	lications
	of body surface area per week for a total of four weeks, or 500 mg once s apart
Hospital. Patient's disease has responded to the previous rituximab treatand The patient has not received rituximab in the previous 6 month and The patient has experienced a relapse and now requires furth and	er treatment of body surface area per week for a total of four weeks, or 500 mg once
INITIATION – CD20+ low grade or follicular B-cell NHL Re-assessment required after 9 months Prerequisites (tick boxes where appropriate)	
The patient has CD20+ low grade or follicular B-cell NH on To be used for a maximum of 6 treatment cycles To be used for a maximum of 6 treatment cycles To be used for a maximum of 6 treatment cycles	

Signed: Date:

HOSPITAL MEDICINES LIST RESTRICTIONS CHECKLIST

PRESCRIBER	PAT	IENT:
Name:	Nan	ne:
Ward:	NHI	
Rituximab (Bi	Riximyo) - continued	
CONTINUATION Re-assessment Prerequisites (1)	ON – CD20+ low grade or follicular B-cell NHL t required after 24 months (tick boxes where appropriate) Rituximab is to be used for maintenance in CD20+ low grade or foll chemotherapy Patient is intended to receive rituximab maintenance therapy for 2	
INITIATION – M Re-assessment	12 cycles) Membranous nephropathy t required after 6 weeks (tick boxes where appropriate)	
	O Patient has biopsy-proven primary/idiopathic membranous noted. Patient has PLA2 antibodies with no evidence of secondary of the progression to end-stage kidney disermeasures (see Note)	eause, and an eGFR of > 60ml/min/1.73m2
CONTINUATION Re-assessment	The total rituximab dose would not exceed the equivalent of 375mg ON – Membranous nephropathy t required after 6 weeks	/m2 of body surface area per week for a total of 4 weeks
0	(tick boxes where appropriate) Patient was previously treated with rituximab for membranous neph	ropathy*
and	Treatment with rituximab was previously successful, but the contreatment Patient achieved partial response to treatment and requires recommendations.	
and	The total rituximab dose used would not exceed the equivalent of 3	75 mg/m2 of body surface area per week for a total of 4 weeks
b) High risk of pc) Conservative dyslipidaemia	marked with * are unapproved indications. progression to end-stage kidney disease defined as > 5g/day proteing measures include renin-angiotensin system blockade, blood-pressina, and anticoagulation agents unless contraindicated or the patient onse defined as a reduction of proteinuria of at least 50% from base	sure management, dietary sodium and protein restriction, treatment of has experienced intolerable side effects.

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PRESCRIBER	PATIENT:	
Name:	Name:	
ard: NHI:		
Rituximab (Riximyo) - continued		
INITIATION – B-cell acute lymphoblastic leukaemia/lymphoma* Re-assessment required after 2 years Prerequisites (tick boxes where appropriate)		
Patient has newly diagnosed B-cell acute lymphoblastic le and Treatment must be in combination with an intensive chemo and The total rituximab dose would not exceed the equivalent of the combination with a second combination with an intensive chemo and	otherapy protocol with curative intent	
Note: Indications marked with * are unapproved indications.		
INITIATION – desensitisation prior to transplant Re-assessment required after 6 weeks Prerequisites (tick boxes where appropriate)		
Patient requires desensitisation prior to mismatched alloge and Patient would receive no more than two doses at 375 mg/r Note: Indications marked with * are unapproved indications.		
INITIATION – pemiphigus* Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) Prescribed by, or recommended by a dermatologist or relevant s by the Health NZ Hospital.	pecialist, or in accordance with a protocol or guideline that has been endorsed	
Patient has severe rapidly progressive pemphigus and Is used in combination with systemic corticosteroids and Skin involvement is at least 5% body surface a or Significant mucosal involvement (10 or more r or Involvement of two or more mucosal sites or Patient has pemphigus and		
Note: Indications marked with * are unapproved indications.		

I confirm that the above details are correct:

Signed: Date:

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.

PRESCRIBER	PATIENT:
Name:	Name:
Ward:	NHI:
Rituximab (Riximyo) - continued	
CONTINUATION – pemiphigus* Re-assessment required after 6 months Prerequisites (tick boxes where appropriate) Prescribed by, or recommended by a dermatologist or relevant special by the Health NZ Hospital.	alist, or in accordance with a protocol or guideline that has been endorsed
Patient has experienced adequate clinical benefit from rituximal ulceration and reduction in corticosteroid requirement and Patient has not received rituximab in the previous 6 months Note: Indications marked with * are unapproved indications.	b treatment, with improvement in symptoms and healing of skin
INITIATION – immunoglobulin G4-related disease (IgG4-RD*) Re-assessment required after 6 weeks Prerequisites (tick boxes where appropriate)	
lowering corticosteroid dose below 5 mg per day (predni	anti-rheumatic drugs is contraindicated or associated with evidence of
Note: Indications marked with * are unapproved indications.	
CONTINUATION – immunoglobulin G4-related disease (IgG4-RD*) Re-assessment required after 12 months Prerequisites (tick boxes where appropriate)	
Treatment with rituximab for IgG4-RD* was previously subut the condition has relapsed Patient is receiving maintenance treatment for IgG4-RD* and Rituximab re-treatment not to be given within 6 months of previously subut the condition has relapsed And Maximum of two 1000 mg infusions of rituximab given two wee	ious course of treatment
Note: Indications marked with * are unapproved indications.	no apart
and analysis management	