## SPECIALISED COMMISSIONING - CLINICAL EVIDENCE EVALUATION CRITERIA FOR CLINICAL COMMISSIONING POLICY PROPOSITION

URN: 1748

TITLE: Addition of rituximab to standard chemotherapy for newly diagnosed CD20

positive-B-cell precursor acute lymphoblastic leukaemia

CRG: Chemotherapy NPOC: Cancer

Lead: Dr Tobias Menne Date: 21 November 2018

This policy is being	For routine	Χ	Not for routine	
considered for:	commissioning		commissioning	
Is the population described in the policy similar to that in the evidence reviewed, including subgroups?	Yes, note the Policy Working Group were not able to identity a particular sub group of adults most likely to benefit. Clinical Panel noted the removal or children from routine commissioning recommendation as had been advised. Panel noted that treatment protocols appear to differ for older adults compared to younger adults and that rituximab may offer greater benefit in younger compared to older adults but accepted the limitation of the evidence base.			
Is the intervention described in the policy similar to the intervention for which evidence is presented in the evidence review?	Yes, the addition of r	ituxima	ab to first line treatment.	
Are the comparators in the evidence reviewed plausible clinical alternatives within the NHS and are they suitable for informing policy development?	Appropriate.			
Are the clinical benefits described in the evidence review likely to apply to the eligible population and/or subgroups in the policy?	although there appearance survival. The macontrolled open-label 209), and one non-ration (n = 282). Panel reco	ared to ain stud phase andomi ognised	survival benefit was shown be some increase in ever dies were one randomised e III trial (GRAALL-2005R; sed open-label phase II trial that the condition is hus that the evidence base	nt n = ial
Are the clinical harms described in the evidence review likely to apply to the eligible and /or ineligible population		rituxir	r, these were not significar mab and control group in t	•

and/or subgroups in the policy?				
The Panel should provide advice on matters relating to the evidence base and policy development and prioritisation. Advice may cover:  Balance between benefits and harms  Quality and uncertainty in the evidence base  Challenges in the clinical interpretation and applicability of policy in clinical practice  Challenges in ensuring policy is applied appropriately  Likely changes in the pathway of care and therapeutic advances that may result in the need for policy review.	Degree of benefit demonstrated appears to be limited and below the threshold that Panel considered sufficient to take forward as a for routine commissioning recommendation.  The policy should progress as a not for routine clinical commissioning policy.			
Overall conclusion	This is a proposition for routine commissioning and	Should proceed for routine commissioning		
	This is a mass soldiers for	Should be reversed and proceed as not for routine commissioning	X	
	This is a proposition for not routine commissioning and	Should proceed for not routine commissioning Should be		
		reconsidered by the PWG		

Report approved by:
David Black
Deputy Medical Director Specialised Services
07 December 2018