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	ER PATIENT:		
Name:			
Nard:	NHI:		
Enteral liqu	quid peptide formula		
INITIATION Prerequisite	I tes (tick boxes where appropriate)		
and	Patient has impaired gastrointestinal function and either cannot tolerate po	lymeric feeds, or polymeric feeds are unsuitable	
	O Severe malabsorption		
	O Short bowel syndrome		
	O Intractable diarrhoea		
	O Biliary atresia or		
	O Cholestatic liver diseases causing malabsorption or		
	O Cystic fibrosis		
	O Proven fat malabsorption or		
0	O Severe intestinal motility disorders causing significant malabsorption or _	Severe intestinal motility disorders causing significant malabsorption	
0	O Intestinal failure		
	The patient is currently receiving funded amino acid formula and		
	The patient is to be trialled on, or transitioned to, an enteral lic	uid peptide formula	
and			
	O A semi-elemental or partially hydrolysed powdered feed has been re	asonably trialled and considered unsuitable	
	For step down from intravenous nutrition		
Note: A reas	asonable trial is defined as a 2-4 week trial.		
CONTINUAT Prerequisite	ATION tes (tick boxes where appropriate)		
	An assessment as to whether the patient can be transitioned to a cows mi formula has been undertaken	k protein or soy infant formula or extensively hydrolysed	
and	The outcome of the assessment is that the patient continues to require an	enteral liquid peptide formula	

I confirm that the above details are correct:	
Cignod	Data