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.

PRESCRIBI	ER		PATIENT:			
Name:			Name:			
Ward:			NHI:			
Elexacaft	or v	with	tezacaftor, ivacaftor and ivacaftor			
INITIATION Prerequisi	_	(tick b	poxes where appropriate)			
and)	Patie	ent has been diagnosed with cystic fibrosis			
and) —	Patie	ent is 6 years of age or older			
		0	Patient has two cystic fibrosis-causing mutations in the cystic fibrosis transmembrane regulator (CFTR) gene (one from each parental allele)			
	or	0	Patient has a sweat chloride value of at least 60 mmol/L by quantitative pilocarpine iontophoresis or by Macroduct sweat collection system			
and						
	or	\circ	Patient has a heterozygous or homozygous F508del mutation			
		0	Patient has a G551D mutation or other mutation responsive in vitro to elexacaftor/tezacaftor/ivacaftor (see note a)			
and (and	O The treatment must be the sole funded CFTR modulator therapy for this condition					
	C	Treat	tment with elexacaftor/tezacaftor/ivacaftor must be given concomitantly with standard therapy for this condition			
Note:						
a) Eligible https://n	mut ctr-c	ations crs.fda	s are listed in the Food and Drug Administration (FDA) Trikafta prescribing information a.gov/fdalabel/services/spl/set-ids/f354423a-85c2-41c3-a9db-0f3aee135d8d/spl-doc			

I confirm that the above details are correct:

C:	D-1	
Signed.	Date:	
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