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		custom synthesised chemicals	d	Commercially available chemicals					
	1 A chemical made in-house specifically to a drug substance manufacturer's requirement	2 A chemical externally made specifically to a drug substance manufacturer's requirement	3 A chemical available for purchase but where the only use is for pharmaceutical manufacture	4 A chemical that is sold as a commodity in a pre-existing, non-pharmaceutical market in addition to its proposed use as a starting material					
Note from Q11 Section 5.2.1:	An applicant should each proposed start of the general princ materials in Section	provide a justifica ting material is app iples for the select 5.1.1	tion for how propriate in light ion of starting	An applicant general not justify the use of commercially availab chemical as a starting	ly need a ble g material.				





This table imps. > th	ation for better health Imp summarizes data and prop ne identification threshold f	urities in API for Case Study 1 osed controls on impurities in the drug substance (e.g., origin for non-mutagenic impurities, potential mutagenic impurities
Impurity	Specified limit in API	Origin (described by applicant to support justification)
i 🤇	Not Specified	Step 6 (specified in G to Not More Than Threshold of Toxicological Concern): a mutagenic impurity; an impurity that persists
ii	0.5%	Step 4b
iii	0.3%	>4 steps upstream of SM A (specified in SM A)
iv	0.3%	Step 5
v	0.3%	>2 steps upstream of SM B (specified in SM B)
vi	0.3%	>2 steps upstream of SM B (specified in SM B)
vii	0.3%	Step 8 (specified in SM G); an enantiomeric impurity; an impurity that persists
viii	0.3%	Step 5
ix	0.2%	Step 5













harmonisation for b npurities in abstance > the in- roposed starting	etter health API: This table su dentification thresho g material or later	ummarizes data on non-mutagenic impurities in the drug old, and mutagenic impurities that may be introduced in a	
Impurity Specified Limit in API		Considerations described by applicant to support justification	
C-9-D1	NMT 1.0%	Impurity introduced in step 4 and is a residual impurity in intermediate C-7, transforms to C-9-D1, a diastereomer of C-9 (the drug substance)	
Individual related substances	NMT 0.10%	All non-mutagenic impurities identified during development of the commercial process	
Mutagenic Impurities		Based on daily dose the M7 threshold for toxicologica concern (TTC) for the API is 25 ppm	
C-8	NMT 25 ppm	Unreacted proposed starting material C-8 from step 6 – known mutagen, spec. set in line with TTC and batch data	
C-6	NMT 25 ppm	Upstream intermediate controlled in an earlier step. However, C-6 impacts the API and should be specified	
C-3, C-4, C-5	Not applicable	C-3, C-4, C-5 do not impact the API, (i.e. they are not specified above 30% the TTC). The applicant chose to control them in intermediate C-7	























