Escalated dose of imatinib for patients with gastro intestinal stromal tumours					
Assessor	initials	s: D	ate:		
Study identifier (Surname of first author + year of publication)					
Type of study Is the study an RCT in which all participants are randomised to imatinib, sunitinib or best supportive care (either provided in addition to imatinib or sunitinib or as only care)? OR Is the study a non-randomised comparative study on patients using either imatinib or sunitinib or best supportive care? OR Is the study case series or case study of more than one patient on same type of diagnosis?		Unclear o next stion	No Exclude		
Participants in the study Does the study contain participants with KIT (CD117) positive unresectable and/or metastatic malignant gastrointestinal stromal tumours (GIST)? ☐ Unresectable ☐ Metastatic Does the study state that disease has progressed on treatment with imatinib at a dose of 400 mg/day? ☐ Yes ☐ No		Unclear	No Exclude		
Does and other comparisons Does the study contain at least one group using escalated doses of imatinib (600mg or 800mg per day)? OR Does the study contain at least one group using sunitinib within its recommended dose range (i.e. 25-75 mg/day)? OR Does the study contain at least one group receiving best supportive care		Unclear O next stion	No Exclude		
Outcomes reported Does the study report any one of the following outcomes? Overall response Overall survival Disease-free survival Progression-free survival Time to treatment failure Health-related quality of life Adverse effects of treatment		Unclear Unclear o next	No Exclude		

Decision	Include	Unclear	Exclude
	Clarification required		