

THORAX



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
ALE.P02.01	II	ALE.P02		sqNSCLC	CLDN1 \geq 50%	Taxanes naive
Astellas 3082-CL-0101	I	3082+Pembrolizumab	G	NSCLC	KRASG12D	Progressed on prior standard therapy Asymptomatic, treated brain metastases
BGB-58067-101	I	BGB-58067	Phase 1b-part A, cohort A	NSCLC	Evidence of homozygous loss of MTAP gene in ctDNA or tumor tissue, or lost MTAP expression in the tumor tissue	2 or 3 lines maximum
		BGB-58067 + Tislelizumab+chemotherapy	Phase 1a – part D dose escalation	NSCLC (non squamous or squamous)	Evidence of homozygous loss of MTAP gene in ctDNA or tumor tissue,	Patient have not received any prior systemic treatment given as primary therapy, or have received 1 cycle of treatment of anti-PD-1/PD-L1 plus platinum-doublet chemotherapy
BI-1479-0009	II	BI 1810631	5	NSCLC	Her2 overexpressed/amplified	
CL1-95035-101	I/II	S095035	1A dose escalation	NSCLC	Homozygous deletion of MTAP	Participants never have already received a MAT2A or PRMT5 inhibitor

THORAX



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
EP0031-101	I/II	EP0031-101	1C	NSCLC	RET fusion-positive NSCLC (prior SRI), monotherapy	Prior selective RET inhibitor
			2A		RET fusion-positive NSCLC, monotherapy	treatment naïve
			2B		RET fusion-positive NSCLC, chemotherapy combination	treatment naïve
GO45416	I/II	GDC-7035 (RO778493)	Expansion phase /A3	NSCLC	KRAS G12D mutation	1 line minimum
GSK-223054	I	GSK5764227	Part 1b dose optimization	ES-SCLC		Radiologically confirmed progression on or after first-line platinum-based chemotherapy for treatment of limited or extensive stage SCLC / Disease progression after PD-1/PD-L1 inhibitor + platinum+ etoposide therapy is required for participants treated with first-line treatment in extensive-stage SCLC
		GSK5764227+ Atezolizumab	Part 1b dose optimization	ES-SCLC		For treatment-naïve participants, no prior systemic therapies in the extensive-stage setting
LOXO-RAS-2001	Ib	LY3537982	B8	NSCLC with brain metastasis	KRAS G12C	Patient must have progressed/be intolerant/ineligible for immunotherapy and platinum based therapy No mutation : EGFR, ALK, BRAF (V600), MET (exon 14), ROS1, RET or NTRK 1/2/3

Senology



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
PanSOHO/BAY2927088	II	BAY 2927088 (per os) : 20mg 2x/j	7	Breast cancer	ER2-	
STX-478-101	I/II	STX-478 + Fulvestrant + Ribociclib	C0	Breast cancer HR+/HER2- or HR+/HER2low measurable disease per RECIST v1.1	PI3Kα H1047X mutations, or other kinase and/or helical domain mutations	At least 1 but no more than 2 prior lines of therapy: <ul style="list-style-type: none"> • CDK4/6 inhibitor, unless the participant is deemed by the investigator intolerant to or ineligible for these agents • Antiestrogen therapy • ≤1 prior line of chemotherapy Or, participants can be treatment-naïve in the metastatic breast cancer setting
		STX-478 + Fulvestrant + Palbociclib	D1	Breast cancer HR+/HER2- or HR+/HER2low measurable disease per RECIST v1.1	PI3Kα H1047X mutations, or other kinase and/or helical domain mutations	Participants must be treatment-naïve in the metastatic breast cancer setting (i.e., no prior systemic therapy for metastatic breast cancer)
		STX-478 + IA ou fulvestrant ou Imlunestrant + Abemaciclib	E0	Breast cancer HR+/HER2- or HR+/HER2low Measurable or non-measurable (bone-only) disease per RECIST v1.1.	PI3Kα H1047X mutations, or other kinase and/or helical domain mutations	At least 1 but no more than 2 prior lines of therapy: <ul style="list-style-type: none"> • CDK4/6 inhibitor, unless the participant is deemed by the investigator intolerant to or ineligible for these agents • Antiestrogen therapy (see inclusion criterion 13) for additional details • ≤1 prior line of chemotherapy Or, participants can be treatment-naïve in the metastatic breast cancer setting
TBBO10203-101	I	TBBO Combo		A breast cancer RH+ HER2-	PIK3CA	

Gynecology



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
CPI-0209	II	CPI0209	M7	Endometrial carcinoma	Food effect cohort ARID1A wild type endometrial carcinoma (up to max 5 patients with concurrent TP53 alterations)	Maximum 2 previous lines including at least one treatment line with systemic platinum-based chemotherapy in advanced/ recurrent disease setting, and anti-programmed cell death protein 1 (PD-1)/ anti-programmed death-ligand 1 (PD-L1) therapy

UROLOGY



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
AGADIR	II	Atezolizumab + BDB001 + radiothérapie	5	UBC	Refractory anti PD-1/L1	
D926UC0001	II	Dato-DXd + Rilvegostomig	6E	Urothelial carcinoma		No prior systemic therapy in the advanced or metastatic setting; Progression >12 months after platinum based neoadjuvant or adjuvant therapy is eligible
DS7300-203	I/Ib	Ifinatamab Deruxtecan (I-DXd)	UC	Urothelial cancer		Relapse or progression after at least 1 prior line of ICI-containing systemic therapy and 1 prior line of systemic chemotherapy, given in combination with other anticancer therapy or separately, with a maximum of 3 prior therapy lines
LOXO-FG3-22001	I	monotherapy	B1	Urothelial cancer		Individual has progressed on or discontinued erdafitinib and Enfortumab vedotin Individuals must also have received all other standard therapies
			B4	Urothelial cancer		Individuals progressed on or discontinued erdafitinib but have not received EV Individuals must also have received all other standard therapies
SNV1521-101	I	SNV1521	Part 3a	Metastatic Castration-resistant Prostate Cancer	Participants harboring known deleterious or suspected deleterious germline or somatic mutations in BRCA1/2, PALB2, and/or RAD51B/C/D by local assay.	May have received up to two lines of cytotoxic chemotherapy in the advanced/metastatic setting. Participants must not have received a prior PARPi

DIGESTIF



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
ART0380C001		ART0380 + irinotécan	B5	CRC	ATM neg	2 lines maximum : - must have previously received fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapy - must NOT have received: Fruquintinib, regorafenib or trifluridine/tipiracil
			B6	pancreas	ATM neg/ATM low	Participant must have mPDAC that has not been previously treated with chemotherapy
BGB-58067-101	I	BGB-58067	Phase Ia – part B, cohort B	PDAC	Evidence of homozygous loss of MTAP gene in ctDNA or tumor tissue, or lost MTAP expression in the tumor tissue.	2 or 3 lines maximum
		BGB-58067 + chemotherapy	Phase Ib – part D	PDAC	Patients with confirmed advanced, metastatic, or unresectable PDAC who have progressed or recurred after standard systemic therapy with MTAP loss	At least 1 line
BI-1479-0009	II	BI1810631	2	Biliary tract / Hepatocellular	Her2 overexpressed/amplified	
			12	CRC	Her2 overexpressed/amplified	
BreAK CRC 001	I	1st line induction COMBO SoC(mFOLFOX6 ± bevacizumab) + STC 1010	Escalation	Stage IIIC, T4b or Stage IV	No BRAFm, MSS/pMMR	Naïf : eligible for 1st line treatment with SOC mFOLFOX6 ± bevacizumab
CL1-95035-101	I	S095035	Escalation 1B	Biliary Tract Cancer	Homozygous deletion of MTAP	Participants never have already received a MAT2A or PRMT5 inhibitor
			Escalation 1C	Pancreatic ductal adenocarcinoma	Homozygous deletion of MTAP	Participants never have already received a MAT2A or PRMT5 inhibitor
MK-3475-158	II	Pembrolizumab	K	Gastric	MSI-high	At least one line
				Small intestine		
				Biliary tract		

SARCOMAS



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
OMX-0407-101	I/II	OMX-0407		Angiosarcoma	Secondary cutaneous AS following radiotherapy, Other cutaneous	At least 1 line and 3lines max

NEUROLOGY

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
MegaMOST	II	Alectinib BID	C	Neuroblastoma	Activating ALK alterations : translocation, mutation	At least one line for metastatic disease, no previous ALK inhibitor (except crizotinib)

SNC

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
F8394-201	II	FORE8394 900mg	Subproto A	SNC +/- metastasis or progressive SNC tumor with BRAF fusion	BRAF fusion (tumor tissue or liquid biopsy)	At least one standard line
			Subproto B	Recurring SNC tumor	BRAF V600E mutation	At least one prior line, radiotherapy included

Solid tumors



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME/EC	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
BGB-43395-101	I/II	BGB-43395	Part A	Breast HR+ or HER2+, prostate, ovarian, endometrial cancer, NSCLC (adenocarcinoma), gastric cancer, esophageal squamous cell carcinoma, colorectal cancer, liposarcoma, squamous cell carcinoma of the head and neck, ewing's sarcoma, familial meningioma, adrenocortical carcinoma	Patients who have previously received standard systemic therapy or for whom treatment is not available or, not tolerated	
BI-1479	II	BI 1810631	6	Agnostic	Her2 overexpressed/amplified	
			10	Agnostic	Her2 mutated	
DSB2455-001	Ia/Ib	DSB2455	Part A : dose escalation	HRD advanced solid tumours, including metastatic castrate-resistant prostate cancer, ovarian cancer and breast cancer	one or more of the following: BRCA1, BRCA2, PALB2, RAD51C, RAD51D	mCRPC: max 1L PARP inhibitor-based Ovarian: at least 1L Platinum-based and 4L in total Breast: max 3L
EZH-1201	I	Tazemetostat	1	Solid Tumors	Moderate hepatic impairment (NCI-ODWG)	At least one line, no prior anti-EZH2
			2	Solid Tumors	Severe hepatic impairment (NCI-ODWG)	At least one line, no prior anti-EZH2

Solid tumors



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME/C	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
F8394-201	II	FORE8394 (900mg) +/-Cobicistat (150mg)	Subproto C	Rare solid tumors (except SNC)	BRAF V600 mutated	Must have received standard therapy Or intolerant to available treatment
			Subproto D	Solid tumors	BRAF alterations	Participants with cutaneous melanoma had previously received and not tolerated a BRAF inhibitor, while participants with thyroid cancer had never received a MAPK inhibitor
GSK-223054	I	GDC-7035 (RO7782493)	Escalation/food effect	Solid tumors	KRAS G12D mutation	One line minimum
IMMUNE 132-15	I	Sacituzumab Govitecan		Subjects with Advanced or Metastatic Solid Tumor and Moderate Liver Impairment	Histologically confirmed advanced or metastatic solid tumor . Creatinine clearance ≥ 30 mL/min, $1.5 \times$ ULN < Total Bilirubin < $3 \times$ ULN	Histologically confirmed advanced or metastatic solid tumor for which no standard therapy is available (TNBC must have received 2 or more prior systemic therapies, including at least 1 for advanced disease)
LOXO-FG3-22001	I	monotherapy	C1	non-urothelial solid tumor	FGFR3 alteration	Participants must be FGFR inhibitor naïve Participants have received all standard therapies

Solid tumors



Pour retrouver tous ces essais et adresser vos patients rapidement : une seule plateforme, www.klineo.com

STUDY NAME	PHASE	MOLECULE	COHORT TYPE			PREVIOUS LINE
			COHORT NUMBER	TUMOR TYPE	SPECIFICITY	
MegaMOST	II	Alectinib BID	C	Solid tumors	Activating ALK alterations : translocation, mutation	At least one line for metastatic disease, no previous ALK inhibitor (except crizotinib)
		Avapritinib	G	Solid tumors	Activating mutations of KIT exon 17 or PDGFRA exon 18 associated or not to mutation on KIT exon 11 or PDGFRA exon 12/14	At least one line for metastatic disease
		Trametinib QD + Dabrafenib BID	F	Solid tumors	BRAF V600 mutation, tumor tissue or liquid biopsy Except melanoma, lung and CRC	
MOST PLUS	II	Nilotinib		PVNS	ABL1, KIT, PDGFRA, PDGFRB, DDR1, DDR2, CSF1R mutations	At least one line
MS201460	I	AntiGD2	I	Sarcomas, glioblastomas resectable		2 prior lines
ODM-212	I/II	ODM-212	Part 2	Malignant pleural mesothelioma (MPM) Epithelioid hemangioendothelioma (EHE) Cholangiocarcinoma (CCA) Head and neck squamous cell carcinoma (HNSCC) Non-small cell lung carcinoma (NSCLC) Colorectal cancer (CRC) Hepatocellular cancer (HCC) Castration-resistant prostate cancer (CRPC) Any other solid tumours with available local data for loss-of-function genetic alterations (truncating mutations or gene deletion) in NF2/LATS1/LATS2 or YAP/TAZ fusions Any other solid tumour based on emerging scientific data as per sponsor's decision		