

### Pancreatic cancer trials open to recruitment – October 2025

This list includes trials targeted at pancreatic cancer alone and trials for many solid tumours which include pancreatic cancer. If you are a patient with pancreatic cancer and wish to discuss whether a clinical trial is right for you, please discuss with your treating Medical Oncologist.

NOTE: Trials listed are available for patients with pancreatic cancer to be referred for consideration. Several trials may have issues with slot availability or waitlists.

#### **Descriptive stages of pancreatic cancer**

Medical Term	What it means
Resectable	This is an early stage of pancreatic cancer where the cancer can still be removed by surgery
Unresectable or locally advanced	The pancreatic cancer has not spread around the body but unfortunately the cancer cannot be removed by surgery due to the tumour size, location, or involvement of surrounding structures
Metastatic	The pancreatic cancer has spread to other parts of the body



# PANCREATIC CANCER TRIALS IN VICTORIA (Trials with specific cohorts for pancreatic cancer)

	(Trials with specific conorts for pancreatic cancer)							
Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)			
NeoFOL-R  Efficacy of Neoadjuvant	N/A	RESECTABLE  Exclusion:	mFOLFIRINOX chemotherapy  Arm A: Surgery then 12 cycles of	Epworth Health	EHJreissatiCentre@epworth.org.au			
FOLFIRINOX in Resectable pancreatic cancer: An international multicentre Randomized, controlled trial (NeoFOL-R) - Australian protocol		Borderline resectable Locally advanced Previous treatment for pancreatic cancer	Arm B: 6 cycles of chemotherapy before surgery followed by 6 of cycles after surgery	Monash Health	gi.oncresearch@monashhealth.org			
,,			Further information:  NeoFOL-R - Victorian Cancer  Trials Link	Alfred Health	act-m@alfred.org.au			
				Western Health	CancerClinicalTrials@wh.org.au			
Investigation of the safety and efficacy of irreversible electroporation (IRE) using the NanoKnife® System in patients	N/A	UNRESECTABLE (Stage 3)  Inclusion: 3 months of chemotherapy (FOLFIRINOX or Gemcitabine based – 1 line only)	NanoKnife System for Irreversible Electroporation (IRE)  IRE is a technique using non- thermal energy to create permanent nanopores in the cell	Peter MacCallum Cancer Centre	SurgicalResearchteam@petermac.or g			
with unresectable stage 3 pancreatic cancer who have received 3 months of chemotherapy			membrane in order to disrupt cellular homeostasis.  Further information:  DIRECT/InspIRE Australia -	Alfred Health	charles.pilgrim@monash.edu			
			Victorian Cancer Trials Link	Epworth Health	EHJreissatiCentre@epworth.org.au			



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AMG193 20230223  A Phase 1b Study Evaluating the	MTAP deletion	METASTATIC/LOCALLY ADVANCED (First line)	AMG 193  AMG 193 is a PMRT5 inhibitor	Peter MacCallum Cancer Centre	PCCTU.MoncC@petermac.org			
Safety, Tolerability, Pharmacokinetics, and Efficacy of		Inclusion:	which is administered orally.  AMG 193 will be administered					
AMG 193 in Combination with Other Therapies in Subjects with		Pancreatic cancer Homozygous MTAP-deletion	with chemotherapy.	Austin Health	samantha.chakar@austin.org.au			
Advanced Gastrointestinal, Biliary Tract, or Pancreatic Cancers with		Exclusion:	Further information: https://trials.cancervic.org.au/det					
Homozygous MTAP-deletion AMG20230223		Prior MAT2A inhibitor or PRMT5 inhibitor	ails/vctl_nct06360354	Epworth Health	EHjreissaticentre@epworth.org.au			
<u>Clarity-PT01</u>	CLDN18.2	METASTATIC/LOCALLY ADVANCED	AZD0901 (antibody-drug conjugate)	Peter MacCallum Cancer Centre	PCCTU.MoncC@petermac.org			
A Phase II, Open-label, Multi- centre Study to Evaluate Safety,		(First line)						
Tolerability, Efficacy, PK, and		Inclusion:	AZD0901 targets and binds to CLDN18.2, which is a protein on					
Immunogenicity of AZD0901 as		Pancreatic cancer	tumour cells, and then delivers					
Monotherapy and in Combination With Anti-cancer Agents in Participants With Advanced Solid		Treatment naïve CLDN18.2 positive	cytotoxic agents which damage these cancer cells.					
Tumours Expressing Claudin 18.2.		Exclusion:	Further information:					
		Exposure to prior CLDN18.2 targeted agents except anti- CLDN18.2 monoclonal antibody	https://trials.cancervic.org.au/det ails/vctl nct06219941					
AMPLICITY (AMP945-202)	FAK	METASTATIC/LOCALLY ADVANCED	<u>Narmafotinib</u>	Epworth	EH-			
A Phase 1b/2a, Multicenter, Open Label Study of the Safety, Efficacy		(First line)	Narmafotinib is an oral FAK inhibitor. Narmafotinib will be		PancreaticCentre@epworth.org.a			
and Pharmacokinetics of Narmafotinib in Combination With Modified FOLFIRINOX in		Inclusion: Treatment naïve for metastatic disease	administered in combination with mFOLFIRINOX chemotherapy		<u>u</u>			
Pancreatic Cancer Patients			Further information:					



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A Phase 1/2, Multicentre, Openlabel, Non-randomized Study to Investigate Safety and Tolerability, Pharmacokinetics, Dosimetry, and Preliminary Activity of 177Lu-FAP-2286 in Patients With an Advanced Solid Tumour	FAP	METASTATIC/LOCALLY ADVANCED (First line) *note second line cohort closed  Inclusion: Pancreatic ductal adenocarcinoma + other cancers  Monotherapy Cohort: Received at least 1 but no more than 2 lines of chemotherapy  Combination cohort: Chemotherapy naïve for advanced disease  Exclusion: Active CNS disease	[177Lu]Lu-FAP-2286  [177Lu]Lu-FAP-2286 is a radiopharmaceutical that targets fibroblast activation protein (FAP). This treatment works by binding to the FAP to allow the targeted delivery of radiation directly to FAP-expressing cancer cells.  [177Lu]Lu-FAP-2286 with or without chemotherapy  Further information:  https://trials.cancervic.org.au/det ails/vctl_nct04939610	Peter MacCallum Cancer Centre	act-m@alfred.org.au  PCCTU.MoncC@petermac.org
Porcupine P2EA  An open-label study to assess the preliminary efficacy and safety of RXC004, (Zamaporvint), in patients with advanced pancreatic cancer who have progressed following therapy with current standard of care.	Porcupine	METASTATIC/LOCALLY ADVANCED (Second line) Inclusion: Progression on first line treatment for incurable disease Mandatory biopsy at enrolment	Zamaporvint  Zamaporvint is an oral porcupine inhibitor. Treatment will be administered with denosumab  Further information: https://www.pancreaticcentre.org.au/treatment/clinical-trials-and-research/porcupine-p2ea	Epworth	EH- PancreaticCentre@epworth.org.au
PAUF-I  A First in Human, Phase 1/2a, Multicentre, Open-label Study Evaluating the Safety, Tolerability,	PAUF	METASTATIC/LOCALLY ADVANCED (Second line +) Inclusion:	PBP1510  PBP1510 is an IgG1 monoclonal antibody (mAb) that targets and neutralises PAUF.	Monash Health	earlyphase.oncresearch@monashhea lth.org



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Pharmacokinetics, and Efficacy of PBP1510 in Patients with Advanced/Metastatic Pancreatic		Received at least 1 line of chemotherapy and progressed	Further information: https://trials.cancervic.org.au/det				
Cancer			ails/vctl nct05141149				
A Phase 1, Open-Label, Multicenter Study of INCB161734 in Participants With Advanced or Metastatic Solid	KRAS G12D	ADVANCED OR METASATIC SOLID CANCERS (Second line +) Inclusion:	INCB161734 INCB161734 in a KRAS G12D inhibitor	Alfred Health	act-m@alfred.org.au		
Tumors With KRAS G12D Mutation		Second line and beyond	Further information: <a href="https://trials.cancervic.org.au/det">https://trials.cancervic.org.au/det</a> <a href="mailto:ails/vctl">ails/vctl</a> nct06179160				
AMG 410	KRAS	ADVANCED OR METASATIC SOLID CANCERS	AMG 410	Peter	PCCTU.EDD@petermac.org		
A Phase 1/1b Study Evaluating the Safety, Tolerability,		(Exhausted standard of care)	AMG 410 is an oral pan-KRAS inhibitor and will be administered	MacCallum			
Pharmacokinetics, Pharmacodynamics, and Efficacy of AMG 410 Alone and in Combination		Inclusion: KRAS mutation or amplification Exhausted conventional	alone or in combination with pembrolizumab and panitumumab	Cancer Centre			
With Other Agents in Participants With KRAS Altered Advanced or Metastatic Solid Tumors		treatment options Pancreatic cohort specified in part 2.	Further information: <a href="https://trials.cancervic.org.au/det">https://trials.cancervic.org.au/det</a> ails/vctl nct07094113				
	HRR deficiency	METASTATIC/LOCALLY ADVANCED	EIK1004	Peninsula and	ag@paso.com.au		
A Phase 1/2, Open-label, Multicenter, Dose-escalation, and Dose-Optimization Study to	uenciency	(Second line +)	EIK1004 is an oral PARP1 inhibitor and will be administered as monotherapy	South Eastern Haematology and			
Evaluate the Safety, Tolerability, and Activity of EIK1004 (IMP1707) as Monotherapy in Participants		Suspected deleterious mutation of select HRR genes	Further information: <a href="https://trials.cancervic.org.au/det">https://trials.cancervic.org.au/det</a>	Oncology Group			
With Advanced Solid Tumors		Exclusion: Prior PARP1 inhibitor	ails/vctl nct06907043				



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ENG19  An open-label, multicenter, Phase I/IIa study assessing the safety and efficacy of EGFR targeted EDVsTM carrying cytotoxic drug PNU-159682 plus concurrent immunomodulatory adjuvant non-targeted EDVs carrying a-galactosyl ceramide in subjects with advanced EGFR-expressing cancers who have failed second-line therapy or where first-and/or second-line therapy is not appropriate (EGFR EDV-D682/GC		METASTATIC/LOCALLY ADVANCED (Third line +)  Inclusion: Progressed on second line or treatment exhausted EGFR expression on local IHC or liquid biopsy	E-EDV-D682/GC  E-EDV-D682/GC is a combination of a EnGelC Dream Vector (EDV) transporting the cytotoxic drug PNU-159682 to cells expressing EGFR and an EDV carrying alphagalactoslyceramide (EDV-GC).  Further information: https://trials.cancervic.org.au/det ails/vctl_actrn12625000203459	Peninsula and South Eastern Haematology and Oncology Group	ag@paso.com.au



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
That Title	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
	(Genomic)	this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
ADCE T03 004	T:	ADVANCED OR METASTATIC	, ,		, , ,
ADCE-T02-001	Tissue Factor (TF)	SOLID CANCERS	<u>AMT-754</u>	Cabrini	clinicaltrials@cabrini.com.au
First-in-Human, Phase 1 Study of			AMT-754 is a targeting tissue		
AMT-754, a Targeting Tissue		Inclusion:	factor antibody-drug conjugate		
Factor Antibody-Drug Conjugate,		Pancreatic cancer	(ADC). ADCs bind to a specific		
in Patients With Advanced Solid		Received ≥1 prior line of therapy	part of the surface of a cancer		
Tumors		No further standard therapy	cell and then deliver targeted	Peninsula and	ag@paso.com.au
		available	treatment directly into the cell.	South Eastern	
				Haematology and	
		Exclusion: Active CNS disease	Further information:	Oncology Group	
			https://trials.cancervic.org.au/de		
	<u> </u>		tails/vctl nct06597721	<u> </u>	
<u>HERTHENA</u>	Her3	ADVANCED OR METASTATIC	HER3-DXd	Monash Health	gi.oncresearch@monashhealth.org
		SOLID CANCERS			
A Study of HER3-DXd in Subjects		Inclusion:	HER3-DXd is a Her3 antibody-		
With Locally Advanced or		Received 1 line of systemic	drug conjugate.		
Metastatic Solid Tumors		therapy and progressed			
		Exclusion:	Further information:		
		Prior anti-Her3 treatment	https://trials.cancervic.org.au/det		
		Prior irinotecan	ails/vctl nct06172478		
			uns/ven needel/21/0		
PRT7732-01	SMARCA4	ADVANCED OR METASTATIC	PRT7732	Monash Health	earlyphase.oncresearch@monashhea
		SOLID CANCERS			<u>Ith.org</u>
A Phase 1 Open-Label, Multi-			PRT7732 is an oral SMARCA2		
Center, Safety and Efficacy Study		Inclusion:	degrader.		
of PRT7732, an Oral SMARCA2		SMARCA4 mutation			
Degrader, in Patients with		Fuelusian	Further information:		
Advanced or Metastatic Solid Tumors with a SMARCA4		Exclusion:	A Study of PRT7732, an Oral		
Nutation		Concomitant SMARCA2 mutation	SMARCA2 Degrader, in Patients		
widtation			with Advanced or Metastatic Solid Tumors with a SMARCA4		
			Mutation - Rare Cancers Australia		
		<u> </u>	ividiation - Nate Cancers Australia		



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	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
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HM-EZHI	SWI/SNF	ADVANCED OR METASTATIC	<u>HM97662</u>	Monash Health	earlyphase.oncresearch@monashhea
	Complex	SOLID CANCERS			
A Phase I, Open-Label,	aberration		HM97662 is an oral medication		<u>Ith.org</u>
Multicenter, Dose Escalation and	(ARID1A	Inclusion:	(EZH1/2 dual inhibitor).		
Expansion Study of HM97662 as a	SMARCA4	SW/SNF Complex aberration		Peninsula and	ag@paso.com.au
Single Agent in Patients With	SMARCA2)	(ARID1A, SMARCA4, SMARCA2)		Southeast	
Advanced or Metastatic Solid			Further information:	Oncology	
Tumors		Exclusion:	https://trials.cancervic.org.au/de	Grampians Health	clinicaltrials@gh.org.au
		Prior valemetostat or other	tails/feed-cta-trial541	(Ballarat)	<u>omnoutriaise griiorgiau</u>
		EZH1/2 inhibitor use		(	
BGB 58067	MTAP loss	ADVANCED OR METASTATIC	BGB-58067	Monash Health	earlyphase.oncresearch@monashhea
		SOLID CANCERS			lth.org
A Phase 1a/b Study Investigating			BGB-58067 is an MTA-		
the Safety, Tolerability,		Inclusion:	Cooperative PRMT5 inhibitor.		
Pharmacokinetics,		MTAP loss (pre-screening			
Pharmacodynamics, and		available)			
Preliminary Antitumor Activity of					
BGB-58067, an MTA-Cooperative		Exclusion:	Further information:	(Note: Austin	
PRMT5 Inhibitor in Patients With		Prior treatment with PRMT5 or	NCT06589596 - Victorian Cancer	Health is only	
Advanced Solid Tumors		MAT2A inhibitor	<u>Trials Link</u>	recruiting GBM)	
BGB 53038	KRAS	ADVANCED OR METASTATIC	BGB-53038	Monash Health	earlyphase.oncresearch@monashhea
	mutation	SOLID CANCERS			lth.org
A Phase 1a/1b Study to	or		BGB-53038 is a pan-KRAS		
Investigate the Safety,	amplificati	Inclusion:	inhibitor.		
Tolerability, Pharmacokinetics,	on .	Evidence of KRAS mutation or			
Pharmacodynamics, and	(excluding	wild-type amplification			
Preliminary Antitumor Activity of					



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BGB-53038, a Pan-KRAS Inhibitor, As Monotherapy or in Combinations in Patients with Advanced or Metastatic Solid Tumors with KRAS Mutations or Amplifications	KRAS G12R)	Exclusion: KRAS G12R mutation Prior treatment with other RAS targeting treatment	Further information: NCT06585488 - Victorian Cancer Trials Link	Peter MacCallum Cancer Centre	PCCTU.EDD@petermac.org
YL211-INT-101-01: A Phase 1, Multicenter, Open-Label, First-in- Human Study to Evaluate the Safety, Tolerability, Pharmacokinetics, and Efficacy of YL211 in Patients With Advanced Solid Tumors	CMET	ADVANCED OR METASTATIC SOLID CANCERS  Inclusion: No further standard treatment options available	YL211  YL211 is a C-MET targeted antibody-drug conjugate (ADC) available for all tumour types for dose escalation where there is either a C-met aberration or where there is a biological rationale for C-met directed therapy.  Further information:  YL211-INT-101-01 - Victorian Cancer Trials Link	Monash Health	earlyphase.oncresearch@monashhea lth.org
MK-1084  A Phase 1, Open-Label, Multicenter Study to Assess Safety, Tolerability, PK, and Efficacy of MK-1084 as Monotherapy and in Combination With Pembrolizumab in Subjects With KRASG12C Mutant Advanced Solid Tumors	KRAS G12C	ADVANCED OR METASTATIC SOLID CANCERS  Inclusion: KRAS G12C mutation	MK1084  MK1084 is an oral KRAS G12C inhibitor.  Further information:  MK-1084 - Victorian Cancer Trials  Link	Monash Health	earlyphase.oncresearch@monashhea lth.org



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PRIMROSE  A Modular Phase I/IIa, Multicentre, Dose Escalation, and Expansion Study of AZD3470, a MTA Cooperative PRMT5 Inhibitor, as Monotherapy and in Combination With Anticancer Agents in Patients With Advanced/Metastatic Solid Tumours That Are MTAP Deficient CS5001  A Phase I, Dose-Escalation and Dose-Expansion Study to Evaluate the Safety, Tolerability, Pharmacokinetics, and Antitumor Activities of CS5001, an Anti-ROR1 Antibody-Drug Conjugate, Used as A Single Agent and in Combination with Systemic Therapies in Patients with Advanced Solid Tumors and Lymphomas.	MTAP deficient	ADVANCED OR METASTATIC SOLID CANCERS  Inclusion: At least 1 prior line of treatment and exhausted treatment options Archival or baseline tumour sample for MTAP testing  Exclusion: Prior PRMT5 inhibitor  ADVANCED OR METASTATIC SOLID CANCERS  Inclusion: Progression on at least 1 prior line of systemic therapy ECOG 0-1	AZD3470  AZD3470 is a novel, potent and selective second-generation PRT5 inhibitor.  Further information: PRIMROSE - Victorian Cancer Trials Link  CS5001  CS5001  CS5001 in an antibody drug conjugate targeting ROR1  Further information: https://www.pancreaticcentre.or g.au/treatment/clinical-trials-and-research/cs5001	Peter MacCallum Cancer Centre	connie.barlas@epworth.org.au or  EH-  PancreaticCentre@epworth.org.a  u
MarkV  A Phase 1a/1b, First-in-Human, Open Label Study to Assess the Safety, Tolerability, and Pharmacokinetics of PMC-309	VISTA	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Progressed on available prior lines of therapy	PMC-309  PMC-309 is an anti-VISTA monoclonal antibody  PMC-309 will be administered	Grampians Health (Ballarat)	clinicaltrials@gh.org.au



Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with	Treatment + Further Information	Site (Where the study	Contact Details (Email the contact person listed
(Anti-VISTA), as Monotherapy and Combined With Pembrolizumab, in Patients With Advanced or Metastatic Solid Tumors		this stage of pancreatic cancer)  Progressed on PD-1 or PD-L1 inhibitor immunotherapy  Exclusion: Prior anti-VISTA therapy	(What the study involves)  alone or in combination with pembrolizumab  Further information: <a href="https://trials.cancervic.org.au/de">https://trials.cancervic.org.au/de</a> <a href="tails/vctl">tails/vctl</a> nct05957081	is being offered) Cabrini	with any enquiries)  clinicaltrials@cabrini.com.au
AT-0174-001  A Phase I Study to Evaluate the Safety, Tolerability, Pharmacology, and Preliminary Efficacy of AT-0174 in Subjects with Advanced Solid Malignancies	IDO1/TDO 2	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Progressed on available prior lines of therapy	AT-0174  AT-0174 is a novel dual inhibitor of IDO1/TDO2  Further information: https://trials.cancervic.org.au/details/vctl_actrn12623000956606	Grampians Health (Ballarat)  St Vincent's Hospital Melbourne	Clinicaltrials@gh.org.au  OncologyTrialCoordinators@svha. org.au
IMPARP  An Open Label, Signal Seeking, Translational, Phase II Trial of Pamiparib in Combination with Tislelizumab in Patients With Advanced Tumours with Homologous Recombination Repair Defects	HRD deficiency	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Second line and beyond Molecular testing within 12 months Confirmed germline or somatic alteration in homologous recombination related gene	Pamiparib and tiselizumab  Pamiparib is a PARP1/2 inhibitor. Tislelizumab is an anti-PD1 antibody.  Further information: <a href="https://trials.cancervic.org.au/details/feed-cta-trial389">https://trials.cancervic.org.au/details/feed-cta-trial389</a>	Western Health	CancerClinicalTrials@wh.org.au
AKTive-001  A Phase 1/1b Multiple Cohort Trial of ALTA2618 in Patients with Advanced Solid Tumors with AKT1	AKT E17K mutation	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Exhausted standard of care	ALTA2618  ALTA2618 is an oral AKT E17K inhibitor	Cabrini	clinicaltrials@cabrini.com.au



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E17K Mutation		therapies  Exclusion: Prior mTOR or PI3K inhibitors Presence of KRAS, NRAS, HRAS or BRAF genomic alterations	Further information: <a href="https://trials.cancervic.org.au/det-ails/vctl-nct06533059">https://trials.cancervic.org.au/det-ails/vctl-nct06533059</a>		
AMT-676-01 First-in-Human, Phase 1 Study of	CDH17	ADVANCED OR METASATIC SOLID CANCERS	AMT-676 AMT-676 is an anti-CDH17	Cabrini	clinicaltrials@cabrini.com.au
AMT-676, an Anti-CDH17 Antibody- Drug Conjugate, in Patients with Advanced Solid Tumors		Inclusion: Second or later line therapy	antibody drug conjugate.  Further information: <a href="https://trials.cancervic.org.au/det_ails/vctl_nct06400485">https://trials.cancervic.org.au/det_ails/vctl_nct06400485</a>		
A Phase 1, Open Label, Dose Escalation and Dose Expansion Study Evaluating the Safety,	KRAS G12C	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: KRAS G12C mutation	D3S 001  D3S 001 is a KRAS G12C inhibitor  Further information:	Cabrini	clinicaltrials@cabrini.com.au
Pharmacokinetics, Pharmacodynamics, and Preliminary Efficacy of D3S 001 Monotherapy in Subjects with Advanced Solid Tumors with a KRAS D.G12C Mutation		*Note pancreatic specific cohort has closed. Pantumour cohort remains open with limited slots	https://trials.cancervic.org.au/det ails/feed-cta-trial449	Peter MacCallum Cancer Centre	PCCTU.EDD@petermac.org
A Phase 1, Open-label, Multicenter Clinical Trial of S095035 (MAT2A Inhibitor) in Adult Participants With Advanced or Metastatic Solid Tumors With Homozygous Deletion	MAT2A MTAP	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Homozygous deletion of MTAP Second line and beyond	S095035  S095035 is an oral MAT2A inhibitor.  Further information:	Alfred Health	act-m@alfred.org.au



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of MTAP			ails/vctl_nct06188702		
KEYNOTE-F49  A Phase 1a/1b, First-in-human, Open-label, Non-randomized, Multicenter, Dose-escalation and Cohort Expansion Study to Evaluate the Safety, Tolerability, Efficacy, Immunogenicity, Pharmacokinetics, and Pharmacodynamics of IOS-1002 Administered Alone and in Combination With a PD-1 Monoclonal Antibody in Advanced Solid Tumors	LILRB1 LILRB2 KIR3DL1 PD-1	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Second or later line	IOS-1002 +/- Pembrolizumab  IOS-1002 binds to LILRB1 (ILT2), LILRB2 (ILT4), and KIR3DL1 receptors on innate and adaptive immune cells that suppress immune responses when activated.  Pembrolizumab in anti-PD1 antibody.  Further information: https://trials.cancervic.org.au/det ails/feed-cta-trial528	Bendigo Health	cancerresearch@bendigohealth.c
A Phase 1/2 First-Time-in-Human, Open-label, Multicenter, Dose Escalation and Expansion Study of the Oral DNA Helicase Werner Inhibitor (WRNi) GSK4418959 Alone or in Combination With Other Anti-cancer Agents in Adult Participants With Mismatch Repair-deficient (dMMR) or Microsatellite Instability-High (MSI-H) Solid Tumors (SYLVER)	WRN dMMR MSI-h	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Second or later line/exhausted treatment options	GSK4418959 +/-PD-1 inhibitor  GSK4418959 is an oral WRN-inhibitor. This is given as monotherapy or in combination with a PD-1 inhibitor.  Further information: https://trials.cancervic.org.au/details/vctl_nct06710847	Peter MacCallum Cancer Centre	PCCTU.MoncB@petermac.org



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
		this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
GDC-7035	KRAS G12D	ADVANCED OR METASATIC SOLID CANCERS	GDC-7035	Peter MacCallum Cancer Centre	PCCTU.EDD@petermac.org
A Phase I/II Dose-Escalation and			GD-7035 is a KRAS G12D		
Expansion Study Evaluating the			inhibitor. Treatment will be		
Safety, Pharmacokinetics, and		Inclusion:	monotherapy or in combination		
Activity of GDC-7035 as a Single		KRAS G12D mutation	with other anti-cancer		
Agent and in Combination With			treatments.		
Other Anti-Cancer Therapies in					
Patients With Advanced Solid			Further information:		
Tumors With a KRAS G12D			https://trials.cancervic.org.au/det		
Mutation			ails/vctl nct06619587		
RO7566802	ανβ8 integrin	ADVANCED OR METASATIC SOLID CANCERS	RO7566802	Peter MacCallum Cancer Centre	PCCTU.EDD@petermac.org
A Phase I, Open-Label,	J		RO7566802 is a αvβ8 integrin		
Multicenter, Dose-Escalation			inhibitor delivered intravenously.		
Study Evaluating the Safety,		Inclusion:	,		
Pharmacokinetics, and Activity of		Second or later line	Further information:		
RO7566802 as a Single Agent and			https://trials.cancervic.org.au/det		
in Combination With			ails/vctl nct06031441		
Atezolizumab in Patients With					
Locally Advanced or Metastatic					
Solid Tumors					
MK-6837-001	TROP2	ADVANCED OR METASATIC SOLID CANCERS	MK-6837 +/- pembrolizumab	Alfred Health	act-m@alfred.org.au
A Phase 1 Open-label, Multicenter			MK-6837 is a TROP2-MMAE		
Study of MK-6837 as		Exclusion:	antibody-drug conjugate		
Monotherapy and Combination		Uncontrolled HIV, Hepatitis B or C	delivered as monotherapy or in		
Therapy in Participants With			combination with PD-1 inhibitor		
Advanced/Metastatic Solid			pembrolizumab.		
Tumors					
			Further information:		
			https://trials.cancervic.org.au/det		
			ails/vctl nct06460961		



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
		this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
<u>BG-C477</u>	CEACAM5	ADVANCED OR METASATIC SOLID CANCERS	<u>BG-C477</u>	Alfred Health	act-m@alfred.org.au
A Multicenter, Open-Label, Phase			BG-C477 is an antibody-drug		
1a/b First-in-Human Study to		Inclusion:	conjugate targeting CEACAM5.		
Investigate the Safety,		CEA >5	BG-C477 will be delivered as		
Tolerability, Pharmacokinetics,			monotherapy or in combination		
Pharmacodynamics, and			with capecitabine and		
Preliminary Antitumor Activity of			bevacizumab.		
BG-C477 in Patients With					
Selected Advanced Solid Tumors			Further information:		
			https://trials.cancervic.org.au/det		
			ails/vctl_nct06596473		
AMT-562-01	HER3	ADVANCED OR METASATIC	AMT-562	Cabrini	clinicaltrials@cabrini.com.au
		SOLID CANCERS			
First-in-Human, Phase 1 Study of			AMT-562 is a novel HER3		
AMT-562, an Anti HER3 Antibody-		Inclusion:	targeting antibody drug		
Drug Conjugate, in Patients with		Second line and beyond	conjugate.		
Advanced Solid Tumors					
			Further information:		
			https://trials.cancervic.org.au/det		
			ails/vctl_nct06199908		
<u>SNT1521</u>	PARP	ADVANCED OR METASATIC SOLID CANCERS	<u>SNT1521</u>	Monash Health	earlyphase.oncresearch@monash
A Phase 1, Open-Label Dose			SNT1521 is a PARP1 inhibitor.		health.org
Escalation and Expansion Study of		Inclusion:			
SNV1521 in Participants With		Exhausted treatment options			
Advanced Solid Tumors			Further information:		
			https://trials.cancervic.org.au/det		
			ails/vctl_nct06220864		
<u>IKSUDA</u>	HER2	ADVANCED OR METASATIC	<u>IKS014</u>	Peninsula and	ag@paso.com.au
		SOLID CANCERS		Southeast	
A Phase 1 Dose Escalation Trial to			IKS014 is a HER2 targeting	Oncology	
Determine the Safety, Tolerance,		Inclusion:	antibody drug conjugate.		



Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)
Maximum Tolerated Dose, and Preliminary Antineoplastic Activity of IKS014, a HER2- Targeting Antibody Drug Conjugate (ADC), in Participants With Advanced HER2+ Solid Tumors		HER2 IHC 1-3+ HER2 ISH negative and positive	Further information: https://trials.cancervic.org.au/det ails/vctl_nct05872295		
LOXO-RAS-200001  A Phase 1a/1b Study of LY3537982 in Patients With KRAS G12C-Mutant Advanced Solid Tumors	KRAS G12C	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: KRAS G12C mutation	LY3537982  LY3537982 is a KRAS G12C inhibitor and will be administered as monotherapy or in combination with other systemic anticancer medications.  Further information: <a href="https://trials.cancervic.org.au/det_ails/vctl_nct04956640">https://trials.cancervic.org.au/det_ails/vctl_nct04956640</a>	Peninsula and Southeast Oncology	ag@paso.com.au
Stingray SR-8541A-001  Phase 1, Dose Escalation, Safety, Tolerability, and Pharmacokinetic Study of SR-8541A (ENPP1 Inhibitor) Administered Orally as Monotherapy in Subjects With Advanced/Metastatic Solid Tumors	ENPP1	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Exhausted conventional treatment	SR-8541A  SR-8541A is an ENPP1 inhibitor.  Further information:     https://trials.cancervic.org.au/det     ails/vctl nct06063681	Peninsula and Southeast Oncology	ag@paso.com.au
Phase 1 Study of a SOS1 Inhibitor, BAY 3498264, in Combination in Participants With Advanced KRASG12C-mutated Solid Tumors	SOS1 KRAS	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: KRAS G12C mutation	BAY3498264  BAY3498264 is a SOS1 inhibitor.  Treatment will be in combination with sotorasib.	Peninsula and Southeast Oncology	ag@paso.com.au



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
	(30,	this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
			Further information:		
			https://trials.cancervic.org.au/det		
			ails/vctl_nct06659341		
AK138D1	HER3	ADVANCED OR METASATIC SOLID	Patritumab Deruxtecan	Peninsula and	ag@paso.com.au
		CANCERS		Southeast	
A First-in-human, Phase I Study of			Patritumab Deruxtecan an anti-	Oncology	
Evaluating Safety, Tolerability,		Inclusion:	HER3 Antibody drug conjugate		
Pharmacokinetics and Preliminary		Exhausted conventional treatment			
Efficacy of AK138D1 in the			Further information:		
Treatment of Advanced Solid			https://trials.cancervic.org.au/det		
Tumors			ails/vctl nct06730386		
DT-7012-CLI-001	CCR8	ADVANCED OR METASATIC SOLID	<u>DT-7012</u>	Peninsula and	ag@paso.com.au
St. 1 ( DT 7042		CANCERS		Southeast	
Study of DT-7012 as a Single			DT 7043 :	Oncology	
Agent and in Combination With an Immune Checkpoint Inhibitor		Inclusion: Second or later line	DT-7012 is an anti-CCR8 antibody.		
in Participants With Advanced		Second or later line	Further information:		
Solid Tumors (DOMISOL)			https://clinicaltrials.gov/study/NC		
John Fumors (Bownson)			T06819735		
SNV4818	PIK3CA	ADVANCED OR METASATIC SOLID	<u>SNV4818</u>	Monash Health	earlyphase.oncresearch@monas
A Disease 4. Ocean Label Dane		CANCERS	CN1/4040 : LDI2/		
A Phase 1, Open-Label Dose Escalation and Expansion Study of		Inclusion:	SNV4818 is an oral PI3Kα		health.org
SNV4818 as Monotherapy or in		Activating PIK3CA mutation	inhibitor. SNV4818 will be delivered with or without		
Combination With Other		Exhausted conventional treatment	fulvestrant		
Anticancer Agents in Participants		Zanadsted conventional treatment	Taivesti aiit		
With Advanced Solid Tumors		Note: Pantumour for dose	Further information:		
		escalation cohorts	NCT06736704 - Victorian Cancer		
CS2009	dMMR	ADVANCED OR METASATIC SOLID	CS2009	Monash Health	earlyphase.oncresearch@monas
	TMB-H	CANCERS			
A Phase I, Dose-Escalation and	HRD		CS2009 is a Tri-specific Antibody		
Dose-Expansion Study to Evaluate	phenotype	Inclusion:	Targeting PD-1/VEGFA/CTLA-4.		



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
	,	this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
the Safety, Tolerability,		Exhausted conventional treatment	CS2009 will be given IV every 3		health.org
Pharmacokinetics and Antitumor			weeks		
Activities of CS2009, a Tri-specific		Note: PDAC considered but other			
Antibody Targeting PD-		tumour types preferred	Further information:		
1/VEGFA/CTLA-4, in Participants		Molecular profile taken into	CS2009 - Victorian Cancer Trials		
With Advanced Solid Tumors		account (i.e. dMMR, TMB-H)	<u>Link</u>		
<u>BM230</u>	HER2	ADVANCED OR METASATIC SOLID	BM230	Monash Health	earlyphase.oncresearch@monash
		CANCERS			
A Phase I, Multicenter, Non-			BM230 is delivered		<u>health.org</u>
randomized, Open-label Study to		Inclusion:	subcutaneously each week for 3		
Evaluate the Safety, Tolerability,		HER2 1+ expression	weeks followed by fortnightly		
Pharmacokinetics, and Efficacy of			administration		
BM230 in Patients With Advanced					
Solid Tumors			Further information:		
			BM230 - Victorian Cancer Trials		
BT317		ADVANCED OR METASATIC SOLID	BT317	Monash Health	earlyphase.oncresearch@monash
		CANCERS			
A Phase I, First-in-human, Open-			BT317 is administered		health.org
label, Dose Escalation Study of		Inclusion:	intravenously		
the Safety, Tolerability,		Exhausted conventional treatment			
Pharmacokinetics, and			Further information:		
Immunogenicity of BNT317 in			NCT06750185 - Victorian Cancer		
Patients with Advanced Solid			Trials Link		
Tumors					



Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with	Treatment + Further Information	Site (Where the study	Contact Details (Email the contact person listed
	(,	this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
OZ-001-101  A Phase 1, Open-label, First-in Human Study to Examine the Safety, Tolerability, Pharmacokinetic Profile, and Preliminary Efficacy of OZ-001 when Administered Orally in Adults with Solid Tumours with a Focus on Triple Negative Breast Cancer	STAT3	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Exhausted conventional treatment  Confirmed accepting PDAC for phase 1a	OZ-001  OZ-001 is a small molecule dual inhibitor of the STAT3 and T-type calcium channels  Further information:  https://trials.cancervic.org.au/det ails/vctl_actrn12625000163404	Peninsula and South Eastern Haematology and Oncology Group	ag@paso.com.au
INI-4001-101  An Open-label, Multiple-Ascending Dose, Two-Part Dose Ranging and Cohort Expansion Study of INI-4001 in Patients with Advanced Solid Tumours	TLR7/8	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Exhausted conventional treatment Pre-screening component (review of medical history)  *Minimal slots available	INI-4001 INI-4001 is TLR7/8 agonist.  Further information: https://trials.cancervic.org.au/det ails/vctl nct06302426	Cabrini	clinicaltrials@cabrini.com.au
LM350-01-10  A Phase I/II, First-in-Human (FIH), Open-Label, Multiple Centre Clinical Study to Evaluate the Safety, Tolerability, Pharmacokinetics, Immunogenicity and Preliminary Efficacy of LM-350 in Patients with Advanced Solid Tumors	CDH17	ADVANCED OR METASATIC SOLID CANCERS  Inclusion: Exhausted conventional treatment	LM350  LM350 is a CDH17 targeted antibody drug conjugate  Further information: https://clinicaltrials.gov/study/NC T07112222?aggFilters=status:not	Peninsula and South Eastern Haematology and Oncology Group	ag@paso.com.au



Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with	Treatment + Further Information	Site (Where the study	Contact Details (Email the contact person listed
		this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
<u>RO7673396</u>	RAS	ADVANCED OR METASATIC SOLID CANCERS	RO7673396	Peter MacCallum Cancer Centre	PCCTU.EDD@petermac.org
A Phase I Dose Escalation and			RO7673396 is an oral RAS		
Expansion Study to Evaluate the		Inclusion:	inhibitor		
Safety, Tolerability,		Exhausted conventional treatment			
Pharmacokinetics and Preliminary		Confirmed presence of RAS	Further information:		
Clinical Activity of RO7673396 as		mutation	https://trials.cancervic.org.au/det		
a Single Agent and in Combination With Other			ails/vctl nct06884618		
Anticancer Therapies in Patients					
With Advanced Solid Tumors					
Harboring RAS Mutation(s)					
AK146D1-102	TROP2 NECTIN4	ADVANCED OR METASATIC SOLID CANCERS	AK146D1	Austin Health	samantha.chakar@austin.org.au
A Phase Ia Clinical Study to			AK146D1 is a an anti-		
Evaluate the Safety, Tolerability,		Exclusion:	TROP2/NECTIN4 bispecific		
Pharmacokinetics and Anti-tumor		Prior TROP2 or NECTIN4 targetting	antibody drug conjugate		
Efficacy of AK146D1 for Injection,		treatments			
an Anti-Trop2/Nectin4 Bispecific		Prior topoisomerase 1 inhibitor	Further information:		
Antibody-drug Conjugate, in		treatment	https://trials.cancervic.org.au/det		
Patients With Advanced Solid Tumors			ails/vctl_nct06929663		



SUPPORTIVE CARE TRIALS FOR PATIENTS WITH PANCREATIC CANCER IN VICTORIA						
Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)	
DINE-PC	N/A	ALL STAGES OF PANCREATIC	Dietetics counselling	Epworth	EH-	
		CANCER			PancreaticCentre@epworth.org.au	
Dietetics Intervention and			Further information:			
Nutritional Evaluation in			https://www.pancreaticcentr			
Pancreatic Cancer Care (DINE-PC)			e.org.au/treatment/clinical-			
			trials-and-research/dine-pc			



### Phase 1 Trials are offered in Victoria at the following locations:

Study Site	Contact	Email	Phone
Alfred Hospital (Prahran VIC 3004)	Clinical Trial Team	$\frac{moncACT1@alfredhealthconnect.onmicrosoft.co}{\underline{m}}$	ТВА
Austin Hospital (Heidelberg VIC 3084)	Samantha Chakar	samantha.chakar@austin.org.au	03 9496 3088
Barwon Health (Geelong VIC 3220)	Karen Aitken	cstu.manager@barwonhealth.org.au	03 4215 2758
Cabrini Malvern (Malvern VIC 3144)	Rochelle Woods	rwoods@cabrini.com.au	95083437
Epworth Hospital (Richmond VIC 3121)	Clinical Research Coordinator	ehjreissaticentre@epworth.org.au	0448 842 680 or 03 9426 8880
PASO Medical (Frankston VIC 3199)	Albert Goikman Clinical Trials Manager	ag@paso.com.au	03 91131307
Monash Health (Clayton VIC 3168)	Early Phase Research Study Coordinator	earlyphase.oncresearch@monashhealth.org	0474 769 510
Peter MacCallum Cancer Centre (Parkville VIC 3052)	Enquiries Line Coordinator	clinicaltrials.enquiries@petermac.org	03 8559 7456 (9am-2pm, Mon-Fri)
St Vincent's Hospital (Fitzroy VIC 3065)	Nadia Ranieri	oncology.research@svha.org.au	03 9231 3167
Western Health (St Albans VIC 3021)	Heike Raunow	CancerClinicalTrials@wh.org.au	03 83959136 or 0434915739

For patients not eligible for any of the above trials, and who have pancreatic cancer that has progressed, are intolerant or are ineligible for standard of care treatments, please consider a phase 1 trial. Clinicians can email or phone the contact listed with any enquiries. Patients, please contact your treating Medical Oncologist if you wish to discuss phase 1 trials.



Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)
CA233-0000/ BMS-986484  A Study of BMS-986484 Alone and Combination Therapy in	CD40/FAP	ADVANCED OR METASTATIC SOLID CANCERS  Exclusion: History of ILD	BMS-986484 (a CD40/FAP bispecific agonist) is delivered as monotherapy or in combination with nivolumab	St Vincent's Hospital Darlinghurst (NSW)	svhs.research@svha.org.au
Participants With Advanced Solid Tumors			with hivolumab	Lyell McEwin Hospital (SA)	Health.NALHNCancerResearch@sa.g  ov.au
Clarity-PT01  A Phase II, Open-label, Multicentre Study to Evaluate Safety, Tolerability, Efficacy, PK, and Immunogenicity of AZD0901 as Monotherapy and in Combination With Anti-cancer Agents in Participants With Advanced Solid Tumours Expressing Claudin 18.2.	CLDN18.2	METASTATIC/LOCALLY ADVANCED  Inclusion: Pancreatic cancer Treatment naïve (first line) CLDN18.2 positive  Exclusion: Exposure to prior CLDN18.2 targeted agents except anti- CLDN18.2 monoclonal antibody	AZD0901 (antibody-drug conjugate)  AZD0901 targets and binds to CLDN18.2, which is a protein on tumour cells, and then delivers cytotoxic agents which damage these cancer cells.  Further information: <a href="https://www.anzctr.org.au/TrialSearch.aspx#&amp;&amp;searchTxt=NCT06219941">https://www.anzctr.org.au/TrialSearch.aspx#&amp;&amp;searchTxt=NCT06219941</a>	Prince of Wales Hospital (NSW)  Fiona Stanley Hospital (WA)	SESLHD-POW- CTRUreferrals@health.nsw.gov.au  Chia.Tan@health.wa.gov.au
AMG193 20230223  A Phase 1b Study Evaluating the Safety, Tolerability, Pharmacokinetics, and Efficacy of AMG 193 in Combination with	MTAP deletion	METASTATIC/LOCALLY ADVANCED  Inclusion: Pancreatic cancer Homozygous MTAP-deletion	AMG 193  AMG 193 is a PMRT5 inhibitor which is administered orally.  AMG 193 will be administered	Genesis Care Norths Shore (NSW)	admin.northshore@genesiscare.com



Trial Title	Tavasta	Torget negulation	Treatment + Further	Site	Contact Details
Trial Little	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Information (What the study involves)	(Where the study is being offered)	(Email the contact person listed with any enquiries)
Other Therapies in Subjects with Advanced Gastrointestinal, Biliary Tract, or Pancreatic Cancers with Homozygous MTAP-deletion AMG20230223		Exclusion: Prior MAT2A inhibitor or PRMT5 inhibitor	with chemotherapy.  Further information: https://www.cancer.nsw.gov.au/r esearch-and-data/cancer-clinical- trials-in-nsw/find-a-cancer- clinical-trial-in-nsw/item?r=13988	Chris O'Brien Lifehouse	jasmine.sell@lh.org.au
AMPLICITY (AMP945-202)  A Phase 1b/2a, Multicenter, Open Label Study of the Safety, Efficacy and Pharmacokinetics of Narmafotinib in Combination With Modified FOLFIRINOX in Pancreatic Cancer Patients	FAK	METASTATIC/LOCALLY ADVANCED (First line) Inclusion: Treatment naïve for metastatic disease	Narmafotinib  Narmafotinib is an oral FAK inhibitor. Narmafotinib will be administered in combination with mFOLFIRINOX chemotherapy  Further information:  AMPLICITY Trial	Genesis Care Norths Shore (NSW)	admin.northshore@genesiscare.com
A FIH study to evaluate the safety, tolerability, pharmacokinetics and pharmacodynamics of VVD-130850, as single agent and in combination with checkpoint inhibition, in participants with advanced solid and hematologic tumors.	STAT3	ADVANCED OR METASTATIC SOLID CANCERS  **STUDY ON HOLD**	VVD130850  VVD130850 is a novel STAT3-inhibitor. Treatment will be as a monotherapy or in combination with checkpoint inhibition (pembrolizumab)  Further information: https://clinicaltrials.gov/study/N CT06188208?term=NCT06188208 &rank=1	Central West Cancer Care Centre (Orange Hospital NSW) Blacktown Cancer & Haematology Centre (NSW) Cancer Research	bernadette.sheldon@health.nsw. gov.au  raymond.tangunan@health.nsw.g  ov.au  admin@crsa.au



PANCREATIC OR SOLID CANCER TRIALS OUTSIDE OF VICTORIA					
Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)
				South Australia (SA)	
	,			Gold Coast University	CBDclinicaltrials@health.qld.gov.a
				Hospital (QLD)	
				ICON Cancer Research (South	admin.southbrisbane@icon.team
				Brisbane QLD)	
ALKOVE-1  A Phase 1/2 Study of the Selective Anaplastic Lymphoma Kinase (ALK) Inhibitor NVL-to655 in Patients with Advanced NSCLC and Other Solid Tumors (ALKOVE-to1)	ALK	ADVANCED OR METASTATIC SOLID CANCERS  Inclusion: ALK rearrangement or activating ALK mutation	NVL655  NVL655 (neladalkib) is an oral selective ALK inhibitor.  Further information: <a href="https://www.cancer.nsw.gov.au/research-and-data/cancer-clinical-trials-in-nsw/find-a-cancer-clinical-trial-in-nsw/item?r=13595">https://www.cancer.nsw.gov.au/research-and-data/cancer-clinical-trials-in-nsw/find-a-cancer-clinical-trial-in-nsw/item?r=13595</a>	Royal North Shore Hospital NSW	PI: malinda.itchins@sydney.edu.au  Trial coordinator:  shirley.liang@health.nsw.gov.au



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with	Information	(Where the study	(Email the contact person listed
		this stage of pancreatic cancer)	(What the study involves)	is being offered)	with any enquiries)
AT-0174-001	IDO1/TDO	ADVANCED OR METASATIC	AT-0174	Royal North	PI:
	2	SOLID CANCERS		Shore Hospital	
A Phase I Study to Evaluate the			AT-0174 is a novel dual inhibitor	(NSW)	helen.wheeler@health.nsw.gov.au
Safety, Tolerability,		Inclusion:	of IDO1/TDO2	, - ,	
Pharmacology, and Preliminary		Progressed on available prior			Trial Coordinator:
Efficacy of AT-0174 in Subjects		lines of therapy	Further information:		
with Advanced Solid Malignancies			https://www.anzctr.org.au/Trial		wenqiong.yu@health.nsw.gov.au
BGB-58067	MTAP	ADVANCED OR METASATIC	BGB-58067	Blacktown Cancer	William.dAvigdor@health.nsw.g
	deficiency	SOLID CANCERS		& Haematology	ov.au
A Phase 1a/b Study Investigating	-		BGB-58067 is a PMRT5 inhibitor	Ţ.	<u>ov.au</u>
the Safety, Tolerability,		Inclusion:		Centre	
Pharmacokinetics,		Any treatment line		(NSW)	
Pharmacodynamics, and			Further information:		
Preliminary Antitumor Activity of			https://www.cancer.nsw.gov.au/r		
BGB-58067, an MTA-Cooperative			esearch-and-data/cancer-clinical-		
PRMT5 Inhibitor in Patients With			trials-in-nsw/find-a-cancer-		
Advanced Solid Tumors			clinical-trial-in-nsw/item?r=14130		
GeneScreen 5FU	DPYD	SOLID CANCERS – ALL STAGES	DPYD genotyping	Lake Macquarie	ClinicalTrialsUnit.LMP@ramsayhealth
DDVD Construes suided dees		la aluai a a	Day and the DDVD and the in-	Private Hospital	
DPYD Genotype-guided dose Personalisation for		Inclusion: Intention to treat with	Pre-emptive DPYD genotyping	(NSW)	<u>.com.au</u>
Fluoropyrimidine prescribing in		Fluoropyrimidine (FP) containing	prior to commencing	•	
Cancer		chemotherapy	Fluoropyrimidine chemotherapy		
		·	Further information:	Fiona Stanley	audrey.margery-
		Exclusion:	ANZCTR - Registration	Hospital (WA)	Muir@health.wa.gov.au
		Prior FP containing		nospital (WA)	iviuii@fiealtif.wa.gov.au
		chemotherapy prior to study			
		entry.			



Trial Title	Targets	Target population	Treatment + Further	Site	Contact Details
	(Genomic)	(Trial suitable for patients with this stage of pancreatic cancer)	Information	(Where the study is being offered)	(Email the contact person listed with any enquiries)
INCB161734  A Phase 1, Open-Label, Multicenter Study of INCB161734 in Participants With Advanced or Metastatic Solid Tumors With KRAS G12D Mutation	KRAS G12D	ADVANCED OR METASATIC SOLID CANCERS (Second line +) Inclusion: Second line and beyond	INCB161734  INCB161734 in a KRAS G12D inhibitor  Further information: https://www.cancer.nsw.gov.au/research-and-data/cancer-clinical-trials-in-nsw/find-a-cancer-clinical-trial-trial-in-nsw/item?r=14020	St Vincent's Hospital Darlinghurst (NSW) Chris O'Brien Lifehouse	robert.kent@svha.org.au  sarah.hing@lh.org.au
BAY3713372  A first in human study to evaluate the safety, tolerability, and pharmacokinetics, pharmacodynamics, and preliminary clinical activity of BAY 3713372, a novel 2 <sup>nd</sup> generation PRMT5 inhibitor in participants with MTAP deleted solid tumors	MTAP loss	ADVANCED OR METASATIC SOLID CANCERS  Inclusion Systemic Treatment naive or treatment exposed advanced pancreatic adenocarcinoma Homozygous MTAP deletion	BAY 3713372  BAY 3713372 is a novel 2 <sup>nd</sup> generation PRMT5 inhibitor.  Further information: <a href="https://www.cancer.nsw.gov.au/research-and-data/cancer-clinical-trials-in-nsw/find-a-cancer-clinical-trial-in-nsw/fitem?r=14234">https://www.cancer.nsw.gov.au/research-and-data/cancer-clinical-trials-in-nsw/find-a-cancer-clinical-trial-in-nsw/fitem?r=14234</a>	Chris O'Brien Lifehouse	teresa.nicholls@lh.org.au



#### PANCREATIC CANCER TRIALS IN SINGAPORE

Trial Title	Targets (Genomic)	Target population (Trial suitable for patients with this stage of pancreatic cancer)	Treatment + Further Information (What the study involves)	Site (Where the study is being offered)	Contact Details (Email the contact person listed with any enquiries)
A phase II study of metronomic capecitabine, oxaliplatin and UGT1A1 genotype-directed irinotecan in metastatic pancreatic cancer patients.	NA	METASTATIC/LOCALLY ADVANCED  Inclusion: Systemic Treatment naive or treatment exposed advanced pancreatic adenocarcinoma	Metronomic capecitabine, oxaliplatin and UGT1A1 genotype directed irinotecan  Further information: https://clinicaltrials.gov/study /NCT05929885?term=NCT059 29885&rank=1b	National Cancer Centre Singapore	honey.shwe.sin@nccs.com.sg
A first in human study to evaluate the safety, tolerability, and pharmacokinetics, pharmacodynamics, and preliminary clinical activity of BAY 3713372, a novel 2 <sup>nd</sup> generation PRMT5 inhibitor in participants with MTAP deleted solid tumors	MTAP loss	ADVANCED OR METASATIC SOLID CANCERS  Inclusion Systemic Treatment naive or treatment exposed advanced pancreatic adenocarcinoma Homozygous MTAP deletion	BAY 3713372  BAY 3713372 is a novel 2 <sup>nd</sup> generation PRMT5 inhibitor.  Further information: https://clinicaltrials.gov/study/NCT06914128	National Cancer Centre Singapore	Wang.jue.lynn@nccs.com.sg
A phase I/II Dose-Escalation and expansion study evaluating the safety, pharmacokinetics, and activity of GDC-7035 as a single agent and in combination with other anti-cancer therapies in patients with advanced solid tumors with a KRAS G12D mutation	KRAS G12D	METASTATIC/LOCALLY ADVANCED  Inclusion: Systemic Treatment refractory KRAS G12D pancreatic adenocarcinoma	GDC-7035 GDC-7035 is a KRAS G12D inhibitor	National Cancer Centre Singapore	Ye.xin@nccs.com.sg



#### PANCREATIC CANCER TRIALS IN SINGAPORE **Trial Title** Target population **Contact Details** Targets Treatment + Further Site (Email the contact person listed (Trial suitable for patients with (Where the study (Genomic) Information this stage of pancreatic cancer) is being offered) with any enquiries) (What the study involves) METASTATIC/LOCALLY **PAUF** National Cancer Goh.mui.leng@singhealth.com.sg PAUF-I PBP 1510 **ADVANCED** Centre Singapore A first in human phase I/2A. PBP 1510 is an anti-PAUF Inclusion: multicentre, open label study antibody Systemic Treatment exposed of evaluating the safety. pancreatic adenocarcinoma tolerability, pharmacokinetics, Further information: and efficacy of PBP1510 in https://clinicaltrials.gov/st patients with udy/NCT05141149 advanced/metastatic

pancreatic cancer