PET/CT Request form / Referral



Date: Name: Address: Medicare No:			Additional Patient Information		
		DOB:	 Diabetic Melanoma Known renal Impairment Previous contrast reaction Public Hospital Outpatient 		
Serum Creatini	ine Level:	eGFR:			
Indication					
Diagnose		Restage	🗆 RT Planning		
Stage		Monitor	Clinical Trial		
Other					
PET/CT All PETCT scans include relevant diagnostic CT 🛛 opt out, low dose CTAC only					
		.			
Primary/Su	spected site				
Histopatho	logy				
	logy	w. Please tick which items apply.	PLEASE NOTE: This form is to be presented at time of appointment with any previous films		
	•••	w. Please tick which items apply. Head & Neck			
Me Lung 161523 Solitar	•••		time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma		
Me Lung Diagnosis	dicare rebateable studies are belo ry Pulmomary Nodule -	Head & Neck	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging		
Me Lung 61523 Solitar Diagnosis 61529 NSCLO	dicare rebateable studies are belo ry Pulmomary Nodule -	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging 61646 Sarcoma - Restaging		
Me Lung Diagnosis	dicare rebateable studies are belo ry Pulmomary Nodule -	Head & Neck 61598 Staging 61604 Restaging	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging		
Me Lung 61523 Solitar Diagnosis 61529 NSCLO Brain 61538 Brain	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown	time of appointment with any previous films Sarcoma Gate or Soft Tissue Sarcoma Staging Gatrointestinal Gastrointestinal Gastrointestinal		
Me Lung 61523 Solitar Diagnosis 61529 NSCLO Brain	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown primary - Staging 	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging 61646 Sarcoma - Restaging Gastrointestinal		
Me Lung	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown primary - Staging Breast	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging 61646 Sarcoma - Restaging Gastrointestinal 61541 Colorectal - Restaging 61577 Oesophageal/GOJ - Staging 61647 Gastroenteropancreatic NET		
Me Lung	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging isy - Evaluation	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown primary - Staging Breast 61524 PET Breast - Stage III, Staging 	time of appointment with any previous films Sarcoma 61640 Bone or Soft Tissue Sarcoma Staging 61646 Sarcoma - Restaging Gastrointestinal 61541 Colorectal - Restaging 61577 Oesophageal/GOJ - Staging 61647 Gastroenteropancreatic NET Diagnosis - DOTA Peptide PET		
Me Lung 61523 Solitar Diagnosis 61529 NSCLO Brain 61538 Brain 61559 Epilep 61560 Alzhe	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging psy - Evaluation imer's - Diagnosis	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown primary - Staging Breast 61524 PET Breast - Stage III, Staging 61525 PET Breast - Restaging	time of appointment with any previous films Sarcoma a 61640 Bone or Soft Tissue Sarcoma Staging b 61646 Sarcoma - Restaging Gastrointestinal b 61541 Colorectal - Restaging b 61577 Oesophageal/GOJ - Staging b 61647 Gastroenteropancreatic NET Diagnosis - DOTA Peptide PET Prostate		
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Me Lung 61523 Solitar Diagnosis 61529 NSCLO Brain 61538 Brain 61559 Epilep 61560 Alzhe Lymphoma 61622 First L treatment 61632 Second	dicare rebateable studies are belo ry Pulmomary Nodule - C - Staging - Restaging osy - Evaluation imer's - Diagnosis	Head & Neck 61598 Staging 61604 Restaging 61610 Metastatic SCC unknown primary - Staging Breast 61524 PET Breast - Stage III, Staging 61525 PET Breast - Restaging Melanoma 61553 Restaging Gynaecology	time of appointment with any previous films Sarcoma biological biological b		

Follow-up appointment with Referring Doctor:			
Practitioner's Name:	Internal Use Only	Yes No	
Address:	Pregnant		
	Front Office Check		
	Patient Identification verified		
	Procedure and consent verified		
	Correct side and site verified		
Signature:	Correct patient data and side ma	Correct patient data and side markers	
5	Tech initials:		
Copy to:	Team leader signature:		
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	BRISBANE PET/CT		Ph: 3840 6200			
Level 3, Mater Private Medical Ctre, 293 Vulture Street, South Brisbane			Fax: 3844 6203			
GREENSLOPES PET/CT Greenslopes Private Hospital, Lower Ground Level, Newdegate Street, Greenslopes		email: petqxr@qldxray.com.au	Ph: 3421 0444 Fax: 3727 7333			
	GA PET/CT Basement, Westside Private Hospital, 32 Morrow Street, Taringa		Ph: 3721 5300 Fax: 3721 5380			
	DREW'S HOSPITAL TOOWOOMBA		Ph: 4633 6828			
Building 2, 280 North Street, Toowoomba		email: qxr.petcttoowoomba@qldxray.com.au	Fax: 4633 6814			
	NTOOWOOMBA hven Street, South Toowoomba		Ph: 4659 4000 Fax: 4659 4088			
	PRIVATE HOSPITAL – HYDE PARK Floor, 9-13 Bayswater Road, Hyde Park, Townsville	email: townsville@qldxray.com.au	Ph: 4759 2800 Fax: 4775 6460			
CAIRN	S PET/CT	email: petctcairns@qldxray.com.au	Ph: 4046 7800			
318 Mu	grave Road, Cairns		Fax: 4051 3028			
	MEDICARE	CRITERIA				
61523	Whole body FDG PET study, performed for evaluation of a solitary pulmonary nod biopsy, or for which an attempt at pathological characterisation has failed (R).	lule where the lesion is considered unsuitable for transthoracic fi	ine needle aspiration			
61524 61525	Whole body FDG PET study, performed for the staging of locally advanced (Stage III Whole body FDG PET study, performed for the evaluation of suspected metastatic					
61529	considered suitable for active therapy (R) (Anaes.) Whole body FDG PET study, performed for the staging of proven non-small cell lu FDG PET study of the brain for evaluation of suspected residual or recurrent malig	ing cancer, where curative surgery or radiotherapy is planned (R)). definitive therapy (or			
	during ongoing chemotherapy) in patients who are considered suitable for furthe Whole body FDG PET study, following initial therapy, for the evaluation of suspect	er active therapy (R).				
61553	suitable for active therapy (R). Whole body FDG PET study, following initial therapy, performed for the evaluation					
61559	suitable for active therapy (R). FDG PET study of the brain, performed for the evaluation of refractory epilepsy w	hich is being evaluated for surgery (R).				
61560	FDG PET study of the brain, performed for the diagnosis of Alzheimer's disease, if: specialist, is equivocal; and	: (a) clinical evaluation of the patient by a specialist, or in consult	ation with a			
	(b) the service includes a quantitative comparison of the results of the study with (c) a service to which this item applies has not been performed on the patient in the service is a service to which the study with the service is a service to which the study with the service is a service to which the service is a service is a service to which the service is a service is a service to which the service is a se		ce database; and			
	(d) a service to which item 61402 applies has not been performed on the patient i		zheimer's disease.			
61563	Applicable not more than 3 times per lifetime (R). Whole body PSMA PET study performed for the initial staging of intermediate to high-risk prostate adenocarcinoma, for a previously untreated patient who is					
61564	considered suitable for locoregional therapy with curative intent. Whole body PSMA PET study performed for the restaging of recurrent prostate adenocarcinoma, for a patient who has undergone prior locoregional therapy and is considered suitable for further locoregional therapy to determine appropriate therapeutic pathways and timing of treatment initiation.					
61565	Whole body FDG PET study, following initial therapy, performed for the evaluation considered suitable for active therapy (R).	n of suspected residual, metastatic or recurrent ovarian carcinon				
61571	Whole body FDG PET study, for the further primary staging of patients with histole conventional staging, prior to planned radical radiation therapy or combined mode	ogically proven carcinoma of the uterine cervix, at FIGO stage IB: dality therapy with curative intent (R).	2 or greater by			
61575	Whole body FDG PET study, for the further staging of patients with confirmed loc chemoradiotherapy or pelvic exenteration with curative intent (R).	al recurrence of carcinoma of the uterine cervix considered suita	able for salvage pelvic			
61577	Whole body FDG PET study, performed for the staging of proven oesophageal or		y (R).			
61604	Whole body FDG PET study performed for the staging of biopsy-proven newly dia Whole body FDG PET study performed for the evaluation of patients with suspect active therapy (R).	ed residual head and neck cancer after definitive treatment, and	who are suitable for			
	Whole body FDG PET study performed for the evaluation of metastatic squamous		s (R).			
61612	Whole body FDG PET study for the initial staging of eligible cancer types, for a pat (a) the eligible cancer type is:	ient who is considered suitable for active therapy, if:				
	 (i) a rare or uncommon cancer (less than 12 cases per 100,000 persons per ye (ii) a typically FDG-avid cancer; and 	ear); and				
	(b) there is at least a 10% likelihood that the PET study result will inform a significa					
61620 61622	Whole body FDG PET study for the initial staging of newly diagnosed or previously Whole body FDG PET study to assess response to first line therapy either during tr	/ untreated Hodgkin or non-Hodgkin lymphoma (R). reatment or within three months of completing definitive first lir	e treatment for			
	Hodgkin or non-Hodgkin lymphoma (R). Whole body FDG PET study for restaging following confirmation of recurrence of					
	Whole body FDG PET study to assess response to second-line chemotherapy if ha Hodgkin lymphoma (R).		-			
	Whole body FDG PET study for initial staging of patients with biopsy-proven bone conventional staging to be potentially curable (R).					
61646	Whole body FDG PET study for the evaluation of patients with suspected residual course of definitive therapy to determine suitability for subsequent therapy with o		r) after the initial			
61647	Whole body 68Ga DOTA peptide PET study, if:(a) a gastro entero pancreatic neuro or equivocal conventional imaging; or(b) both:(i) a surgically amenable gastro en	pendocrine tumour is suspected on the basis of biochemical evic				
	conventional techniques; and(ii) the study is for excluding additional disease sites					
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0	r more mormation, please visit quaxiay.com.au/patients/r	esuits-portai/				

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