Indication	For the treatment of metastatic hormone-relapsed prostate cancer before chemotherapy is indicated.					
	NB No previous treatment with enzalutamide or darolutamide or apalutamide or					
	abiraterone should have been received OR if the patient has previously received					
	enzalutamide it was stopped within 3 months of starting due to dose limiting toxicity and					
	there is clear absence of disease progression.					
Treatment	Palliative					
Intent						
Frequency and	Repeat every 28 days					
number of						
cycles	Continue until disease progression, unacceptable toxicity or patient choice.					
Monitoring	Monitor U&Es and FBC with each cycle for 6 months and then every 3 months					
Parameters	thereafter if clinically indicated.					
pre-treatment	• Monitor LFTs every 2 weeks for first 3 months then monthly for 3 months and then every 3 months thereafter if clinically indicated.					
	Blood pressure, serum potassium and fluid retention should be monitored before					
	treatment and at least monthly thereafter. In patients at high risk of congestive heart					
	failure monitoring should be 2 weekly for the first 3 months of treatment, then if					
	clinically stable monthly thereafter.					
	Use with caution if history of cardiovascular disease (before treatment hypertension					
	must be controlled and hypokalaemia corrected, consider maintaining potassium levels					
	at >/=4mmol/L during treatment). New patients with cardiac failure should have an					
	ECHO (transthoracic echocardiogram to measure Left Ventricular Ejection Fraction)					
	before starting treatment.					
	Hepatic impairment: no dose adjustment in pre-existing mild (Child-Pugh A)					
	impairment. Limited data available in moderate (Child-Pugh B) impairment, to be					
	prescribed at clinician's decision. Not to be used in severe hepatic impairment.					
	Renal Impairment: Use with caution in severe renal impairment.					
	Management of adverse reactions and dose adjustments:					
	<ul> <li>Hepatotoxicity and hepatic impairment:</li> <li>If increase in ALT&gt;5x ULN, discontinue treatment and all other concomitant</li> </ul>					
	<ul> <li>If increase in ALT&gt;5x ULN, discontinue treatment and all other concomitant medications that are potentially hepatotoxic. Re-treatment may take place only</li> </ul>					
	after return of liver function tests to the patient's baseline, and at the reduced					
	dose level of 500mg once a day with serum transaminases monitored at least					
	every two weeks for three months and monthly thereafter. No further dose					
	reduction is permitted; if hepatotoxicity recurs at the reduced dose treatment					
	should be discontinued.					
	<ul> <li>If patients develop severe hepatotoxicity (ALT 20 x ULN) discontinue treatment</li> </ul>					
	and do not re-treat.					
	• For patients who develop Grade >/= 3 toxicities including hypertension,					
	hypokalaemia, oedema and other non-mineralocorticoid toxicities, treatment					
	should be withheld and appropriate medical management should be instituted.					
	Treatment should not be reinitiated until symptoms of the toxicity have resolved					
	to Grade 1 or baseline.					
	<ul> <li>Prednisolone dose may be reduced to 5 mg od at clinicians' discretion.</li> </ul>					
	<u>Common drug interactions (for comprehensive list refer to BNF/SPC):</u>					
	• Caution is recommended in patients concomitantly treated with drugs known to be					
	associated with myopathy/rhabdomyolysis (e.g glucocorticoids, cholesterol lowering					
	drugs, zidovudine, amiodarone, colchicine)					

Protocol No	URO-021	Kent and Medway SACT Protocol Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V5	Written by	M.Archer	
Supersedes version	V4	Checked by	C.Waters M.Capomir	
Date	02.08.2022	Authorising consultant (usually NOG Chair)	A.Edwards	

	<ul> <li>Strong inducers (e.g. phenytoin, carbamazepine, rifampicin) of CYP3A should be avoided or used with caution.</li> </ul>
	<ul> <li>Dose reduction of medicines with a narrow therapeutic index metabolized by CYP2D6 (e.g metoprolol, propranolol, venlafaxine, haloperidol, risperidone) should be considered.</li> </ul>
	• Use with caution when given concomitantly with other medications known to prolong QT interval.
	• Avoid spironolactone, co administration with abiraterone is not recommended.
	• Hyperglycaemia/Hypoglycaemia: The risk of hypoglycaemia has been linked to co- administration with pioglitazone or repaglinide. Patients with diabetes should be advised to closely monitor their blood sugars and liaise with their diabetic team. Close monitoring for toxicity is recommended for patients taking CYP2C8 substrate with a narrow therapeutic index (e.g pioglitazone and repaglinide).
	• <b>Missed dose</b> : in the event a dose of abiraterone or prednisolone is missed, the patient
	should not take the dose and wait until the next scheduled dose to continue.
	• For oral self-administration: refer to local Trust policy on oral anti-cancer medicines and supply Patient Information Leaflet.
References	KMCC proforma URO-021 v4 SPC accessed online 02.08.2022 CDF list v1.223 accessed
	online 02.08.2022 BT form accessed online 02.08.2022 Urology NOG 29 March 2022

NB For funding information, refer to CDF and NICE Drugs Funding List

## Repeat every 28 days

TTO	Drug	Dose	Route	Directions
Day 1	ABIRATERONE	1000mg	РО	Each day continuously for 28 days. Tablets should be taken at least 1 hour before food or at least 2 hours after eating. Swallow whole with water. Available as 500mg tablets.
	PREDNISOLONE	5mg	РО	BD for 28 days

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