• • • • • •		alarih fartha trantoret afi
Indication		below by the treatment of: a syndrome with non-locally advanced, non-meteriatic multiple ($\sum (-5)$ BCCs clinically
		n syndrome with non-locally advanced, non-metastatic multiple (>/=6) BCCs clinically
	and	nt lesions at the point of decision to treat of which 3 are at least 5mm
		ocally advanced, non-metastatic multiple BCC (>/=6) clinically evident lesions at the point of
		on to treat of which 3 are at least 5mm AND are appropriate for surgery.
	ueersie	on to treat of which 5 are at least 5mm AND are appropriate for surgery.
	NB: Pa	atients must be appropriate for surgery i.e. surgically eligible tumours.
		rust policy regarding the use of unlicensed treatments must be followed as this is an
		ensed dosing schedule and indication for vismodegib.
Treatment	Palliat	tive
Intent		
Frequency a	nd Vismo	odegib 150mg daily for 12 weeks, followed by 8 weeks off treatment.
number of c		nittent schedule repeated for a total of 72 weeks
Monitoring	• M	Ionitor FBC, LFT's and U&E's prior to each cycle. If neuts <1.0 and/or PLT <100 d/w
Parameters		onsultant.
treatment	• W	/CBP must have a negative pregnancy test, conducted by a health care provider within 7
		ays before starting treatment and have a negative pregnancy test monthly during
	tre	eatment, even if the patient becomes amenorrhoeic.
	• He	epatic impairment: no dose adjustment required.
	• Re	enal impairment: no dose adjustment required in mild and moderate renal impairment.
	Lir	mited data in severe impairment, patients should be monitored closely for adverse
	re	eactions.
	• M	lanagement of adverse reactions and dose adjustments:
	• Do	ose adjustments are not recommended.
	• Se	evere cutaneous adverse reactions (SCARs) including cases of Stevens-Johnson
	-	ndrome/Toxic epidermal necrolysis (SJS/TEN), Drug reaction with eosinophilia and systemic
		mptoms (DRESS) and acute generalised exanthematous pustulosis (AGEP), have been
		eported during treatment with vismodegib, if the patient develops any of these reactions'
		eatment with vismodegib must be discontinued and not restarted at any time.
		ommon drug interactions (for comprehensive list refer to BNF/SPC):
		void concomitant treatment with potent CYP inducers (e.g rifampicin, phenytoin, St. Johns'
		/ort, carbamazepine).
		ismodegib has the potential to act as an inhibitor of breast cancer resistance protein (BCRP)
		nd may give rise to increased exposure of medicinal products transported by this protein,
		uch as rosuvastatin, topotecan, and sulfasalazine. Concomitant administration should be
		erformed with caution and a dose adjustment may be necessary.
		Tissed dose: If a dose is missed the patient should not take an additional dose and take at
		ne dose at the next scheduled time.
		regnancy/contraception
		ismodegib must not be used during pregnancy and female and male patients will be punselled as described below.
		ounselling for female patients:
 I confirm the patient has been counselled about the adverse use of vismodegib in p AND, if a woman of child hearing notantial, has been advised that she should use to 		
		ND, if a woman of child-bearing potential, has been advised that she should use two forms f contraception (including one highly effective method and one barrier) during vismodegib
		nerapy and for 24 months after the final dose, AND has had a negative medically supervised
		regnancy test within the past seven days.
Protocol No	SKI-005	Kent and Medway SACT Protocol
		Disclaimer: No responsibility will be accepted for the accuracy of this information when used else-
Version	V3	where. Written by M.Archer
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		where.		
Version	V3	Written by	M.Archer	
Supersedes	V2	Checked by	C.Waters V3	
version			H.Paddock V2	
			V3 minor change pharmacist approval only	
Date	26.04.2023	Authorising consultant (usually NOG Chair)	J.Turner V2	

	 Counselling for male patients: I confirm that the patient has been counselled about the adverse use of vismodegib in relation to pregnancy and has been advised that he should always use a condom (with spermicide if available), during vismodegib therapy and for 2 months after the final dose. Male patients should not donate semen while taking vismodegib and for 2 months after the final dose. Patients must comply with the "Erivedge Pregnancy Prevention Programme" materials. Blood donation: Patients should not donate blood while taking Erivedge and for 24 months after the final dose. For oral self-administration: refer to local Trust policy on oral anti-cancer medicines and supply Patient Information Leaflet and Macmillan information sheet.
References	Blueteq form accessed online 18.10.21 SPC accessed online 21.10.21 KMCC proforma SKI-005 v1

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

Intermittent schedule of vismodegib given for 12 weeks, followed by 8 weeks off treatment. Repeated for a total of 72 weeks (4 blocks of vismodegib treatment).

TTO	Drug	Dose	Route	Directions
Day 1	VISMODEGIB	150mg	PO	OD for 12 weeks followed by an 8-week break. Swallow whole with water with or without food. Available as 150mg capsule Do not open capsules.
	Metoclopramide	10mg	РО	Up to TDS PRN Do not take for more than 5 days continuously.
	Loperamide	2-4mg	PO	Take 4mg (2 capsules) initially, then 2mg (1 capsule) after each loose stool when required. Maximum 16mg (8 capsules) a day. Dispense 30 capsules on cycle 1 then only if required.

Protocol No	SKI-005	Kent and Medway SACT Protocol		
		Disclaimer: No responsibility will be accepted for the accuracy of this information when used else-		
		where.		
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