## SA2032 - Somatropin

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Reg No:	First Names:	First Names:
Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Somatropin		
Initial application — growth hormone deficience Applications only from a paediatric endocrinologis Prerequisites(tick boxes where appropriate)	y in children t or endocrinologist. Approvals valid for 9 months.	
cardiomyopathy, hepatic dysfunction	ng symptomatic hypoglycaemia, or with other signification) and diagnosed with GH < 5 mcg/l on at least two shed hypoglycaemia (whole blood glucose < 2 mmol	random blood samples in the first 2 weeks of
Height velocity < 25th perce standards of Tanner and Data	ntile for age adjusted for bone age/pubertal status if a vies (1985)	appropriate over 6 or 12 months using the
A current bone age is < 14 y	rears (female patients) or < 16 years (male patients)	
	of < 5.0 mcg per litre in response to two different groing with sex steroid priming is required	owth hormone stimulation tests. In children who
and	ed for a malignancy, they should be disease free for a	at least one year based upon follow-up
	maging appropriate for the malignancy, unless there	
and	ituitary gland has been obtained	
	mutary grand has been obtained	
Renewal — growth hormone deficiency in child	lren	
Current approval Number (if known):		
` ' '	or endocrinologist. Approvals valid for 12 months.	
Prerequisites(tick boxes where appropriate)		
A current bone age is 14 years or and	under (female patients) or 16 years or under (male patients)	atients)
Height velocity is greater than or e	qual to 25th percentile for age (adjusted for bone age over six months using the standards of Tanner and D	
	qual to 2.0 cm per year, as calculated over 6 months	
No serious adverse effect that the	patients specialist considers is likely to be attributabl	e to growth hormone treatment has occurred
No malignancy has developed sind	ce starting growth hormone	
Initial application — Turner syndrome		
	t or endocrinologist. Approvals valid for 9 months.	
The patient has a post-natal genot	ype confirming Turner Syndrome	
	over 6-12 months using the standards of Tanner and	d Davies (1985)
A current bone age is < 14 years		

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Name	:	Surname:	Surname:
Addre	SS:	DOB:	Address:
		Address:	
Fax N	umber:		Fax Number:
Som	atropin - continued		
Rene	ewal — Turner syndrome		
Curre	ent approval Number (if known):		
	cations only from a paediatric endocrinologist equisites(tick boxes where appropriate)	or endocrinologist. Approvals valid for 12 months.	
	Height velocity is greater than or ended Ranke's Turner Syndrome growth wand	qual to 50th percentile for age (while on growth horm velocity charts)	one calculated over 6 to 12 months using the
	Height velocity is greater than or e	qual to 2 cm per year, calculated over six months	
	A current bone age is 14 years or u	under	
	No serious adverse effect that the	specialist considers is likely to be attributable to grow	rth hormone treatment has occurred
	and  No malignancy has developed since	ce starting growth hormone	
App	Il application — short stature without grow ications only from a paediatric endocrinologis equisites(tick boxes where appropriate)	rth hormone deficiency t or endocrinologist. Approvals valid for 9 months.	
	delay	standard deviations below the mean for age or for b	one age if there is marked growth acceleration or
	the standards of Tanner and Davie	for age (adjusted for bone age/pubertal status if app s(1985)	ropriate), as calculated over 6 to 12 months using
	A current bone age is < 14 years o	r under (female patients) or < 16 years (male patient	s)
	The patient does not have severe of medications known to impair heigh	chronic disease (including malignancy or recognized it velocity	severe skeletal dysplasia) and is not receiving
Renewal — short stature without growth hormone deficiency			
	-	•	
	ent approval Number (if known): cations only from a paediatric endocrinologist	or endocrinologist. Approvals valid for 12 months.	
	equisites(tick boxes where appropriate)	о опасотно од от трротато нападот д	
	12 months using the standards of	qual to 50th percentile (adjusted for bone age/pubert Tanner and Davies (1985)	al status if appropriate) as calculated over 6 to
	and	qual to 2 cm per year as calculated over six months	
		qual to 2 on por your as salealated ever six months	
	and	under (female patients) or 16 years or under (male patients)	atients)
	A current bone age is 14 years or the and		

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Reg No:			First Names:	First Names:
Name:			Surname:	Surname:
Addres	s:		DOB:	Address:
			Address:	
Fax Nu	ımber:			Fax Number:
Soma	tropin	- continued		
Appliendoc Prere	cations o	t. Approvals valid for 9 months. (tick boxes where appropriate)  The patient's height is more than 2  Height velocity is < 25th percentile standards of Tanner and Davies (1  A current bone age is to 14 years of the patient is metabolically stable.  The patient is under the supervision  The patient has a GFR less creatinine (umol/l)) × 40 = co	et, endocrinologist or renal physician on the recomme 2 standard deviations below the mean 4 (adjusted for bone age/pubertal status if appropriate	e) as calculated over 6 to 12 months using the alle patients) sence of any other severe chronic disease the Schwartz method (Height(cm)/plasma may not be receiving dialysis
Renev	wal — sl	nort stature due to chronic renal i	nsufficiency	
Applic	ations or rinologis	val Number (if known):nly from a paediatric endocrinologist t. Approvals valid for 12 months. (tick boxes where appropriate)	t, endocrinologist or renal physician on the recommer	ndation of a paediatric endocrinologist or
	and	Height velocity is greater than or e 12 months using the standards of	qual to 50th percentile (adjusted for bone age/pubert Tanner and Davies (1985)	al status if appropriate) as calculated over 6 to
	and	Height velocity is greater than or e	qual to 2 cm per year as calculated over six months	
		A current bone age is 14 years or	under (female patients) or 16 years or under (male pa	atients)
	and	No serious adverse effect that the	patients specialist considers is likely to be attributable	e to growth hormone has occurred
	and	No malignancy has developed after	er growth hormone therapy was commenced	
	and	The patient has not experienced s	ignificant biochemical or metabolic deterioration confi	irmed by diagnostic results
	and		transplantation since starting growth hormone treatm	
	and	If the patient requires transplantati made after transplantation based of	on, growth hormone prescription should cease before on the above criteria	e transplantation and a new application should be

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Address:	DOB:	Address:	
	Address:		
Fax Number:Somatropin - continued		Fax Number:	
and  The patient is aged six months or of and  A current bone age is < 14 years (for and and)  Sleep studies or overnight oximetry obstructive sleep disorder is found surgeon  and  The patient is aged two and  There is no evidence of equal to 0.5 standard of and	der-Willi syndrome that has been confirmed by genet older female patients) or < 16 years (male patients) y have been performed and there is no obstructive sle, it has been adequately treated under the care of a page of the control of the care of a page of the control of the care of a page of the control of the care of a page of the control of the care of a page of the control of the care of a page of the control of the care of a page of the control of the control of the care of a page of the control of the care of a page of the control of the care of the control of the control of the care of the care of the care of the control of the care of the	eep disorder requiring treatment, or if an aediatric respiratory physician and/or ENT  BMI that has increased by greater than or avay assessment is planned to be undertaken	
Renewal — Prader-Willi syndrome			
Current approval Number (if known):			
Applications only from a paediatric endocrinologist <b>Prerequisites</b> (tick boxes where appropriate)	or endocrinologist. Approvals valid for 12 months.		
Height velocity is greater than or ed 12 months using the standards of and	qual to 50th percentile (adjusted for bone age/pubert Tanner and Davies (1985)	al status if appropriate) as calculated over 6 to	
Height velocity is greater than or ea	qual to 2 cm per year as calculated over six months		
	under (female patients) or 16 years or under (male pa	atients)	
	patient's specialist considers is likely to be attributable	e to growth hormone treatment has occurred	
	er growth hormone therapy was commenced		
	e II diabetes or uncontrolled obesity as defined by BM eding 12 months	Il that has increased by greater than or equal to	

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		NOVCITIBET ZUZU
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Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Somatropin - continued		
treatment of a pituitary tumour)  and The patient has undergone appropriated The patient has severe growth horrore and The patient's serum IGF-I is more to and The patient has poor quality of life, growth hormone deficiency (QoL-At	n that is known to cause growth hormone deficiency riate treatment of other hormonal deficiencies and ps mone deficiency (see notes) than 1 standard deviation below the mean for age an as defined by a score of 16 or more using the disease GHDA®)	sychological illnesses  d sex se-specific quality of life questionnaire for adult
equal to 3 mcg per litre during an adequately performance patients with one or more additional anterior pituital isolated growth hormone deficiency require two grown additional test is required, an arginine provocation The dose of somatropin should be started at 0.2 mean normal value for age and sex; and Dose of somatropin not to exceed 0.7 mg per day for the performance of the started at 0.2 mg per day for the performance of t	severe growth hormone deficiency is defined as a permed insulin tolerance test (ITT) or glucagon stimularity hormone deficiencies and a known structural pituitowth hormone stimulation tests, of which, one should on test can be used with a peak serum growth hormone globally and be titrated by 0.1 mg monthly until the second male patients, or 1 mg per day for female patients rism, patients must be monitored for any required adj	tion test. tary lesion only require one test. Patients with be ITT unless otherwise contraindicated. Where one level of less than or equal to 0.4 mcg per litre. erum IGF-I is within 1 standard deviation of the

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APPLICAN	NT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:		First Names:	First Names:
Name:		Surname:	Surname:
Address: .		DOB:	Address:
		Address:	
Fax Numb	er:		Fax Number:
Somatro	pin - continued		
Renewal	<ul> <li>adults and adolescents</li> </ul>		
Applicatio	pproval Number (if known):ns only from a paediatric endocrinologis	st or endocrinologist. Approvals valid for 12 months.	
	and There has been an improve Growth Hormone Deficience and Serum IGF-I levels have be	ed with somatropin for < 12 months  ement in Quality of Life defined as a reduction of at leady in Adults (QoL-AGHDA®) score from baseline  en increased within ±1SD of the mean of the normal resources.	range for age and sex
or  The dose of somatropin has not exceeded 0.7 mg per day for male patients, or 1 mg per day for female patients  The patient has been treated with somatropin for more than 12 months  and			
The patient has not had a deterioration in Quality of Life defined as a 6 point or great score on treatment (other than due to obvious external factors such as external stress and  Serum IGF-I levels have continued to be maintained within ±1SD of the mean of the obvious external factors)  and  The dose of somatropin has not exceeded 0.7 mg per day for male patients or 1 mg		tressors) the normal range for age and sex (other than for	
or	The patient has had a Spectrenewal criteria under this i	cial Authority approval for somatropin for childhood de ndication	ficiency in children and no longer meets the
	The patient has undergone and	appropriate treatment of other hormonal deficiencies	and psychological illnesses
		wth hormone deficiency (see notes)	
		s more than 1 standard deviation below the mean for a	age and sex
		y of life, as defined by a score of 16 or more using the iency (QoL-AGHDA®)	disease-specific quality of life questionnaire for
equal to 3 Patients v isolated g an additio The dose mean nor	mcg per litre during an adequately perf vith one or more additional anterior pitui rowth hormone deficiency require two g nal test is required, an arginine provoca of somatropin should be started at 0.2 mal value for age and sex; and	s, severe growth hormone deficiency is defined as a property formed insulin tolerance test (ITT) or glucagon stimulated the hormone deficiencies and a known structural pitule rowth hormone stimulation tests, of which, one should tion test can be used with a peak serum growth hormore daily and be titrated by 0.1 mg monthly until the second service of the second second service of the second second service of the second s	tion test.  itary lesion only require one test. Patients with I be ITT unless otherwise contraindicated. Where one level of less than or equal to 0.4 mcg per litre. erum IGF-I is within 1 standard deviation of the
	nmencement of treatment for hypopituit	rifor male patients, or 1 mg per day for female patients arism, patients must be monitored for any required adj	