



Gauntlet Brace Compliance Packet

This Packet has 10 Pages to insure proper documentation and DME compliance.

Pages:

- 1) Gauntlet Brace Compliance Packet and Checklist
- 2) Biomechanical Evaluation (page 1)
- 3) Biomechanical Evaluation (page 2)
- 4) Tests and Gait Evaluation
- 5) Document of Medical Necessity Custom Molded Gauntlet
- 6) Prescription (Rx) Custom Molded Gauntlet
- 7) G800 Balance Brace Gauntlet AFO Order Form
- 8) Proof of Delivery
- 9) Medicare DMEPOS Supplier Standards
- 10) Dispensing Documentation Custom Molded Gauntlet

Below is a check list to ensure all items are completed

TO BE COMPLETED BY PHYSICIAN:	
Biomechanical Evaluation for Patient Medical Record	
☐ Medical necessity documents	
Medical Necessity Form	
☐ Supports AFO usage qualification	
☐ Evaluates necessity for prefabrication device	
☐ Demonstrates necessity for custom fitting	
☐ Support for selection code(s)	
Prescription	
☐ Patient name (printed)	
☐ Item Description	
☐ Left ☐ Right ☐ Bilateral	
☐ Diagnosis	
☐ Clinician name (printed)	
☐ Clinician signature	
□ Date	
☑ TO BE GIVEN TO PATIENT:	
Proof of Delivery (provide copy to patient)	
☐ Patient name (printed)	
" '	
☐ Patient address	
" ·	
☐ Patient address	
☐ Patient address ☐ Item description	
☐ Patient address ☐ Item description ☐ Item code(s):	

☑ TO BE COMPLETED BY SUPPLIER/CLINICIAN

Dispensing Chart Notes

- \square Orthosis type
- □ Demonstrates fitting
- ☐ Document patient satisfaction





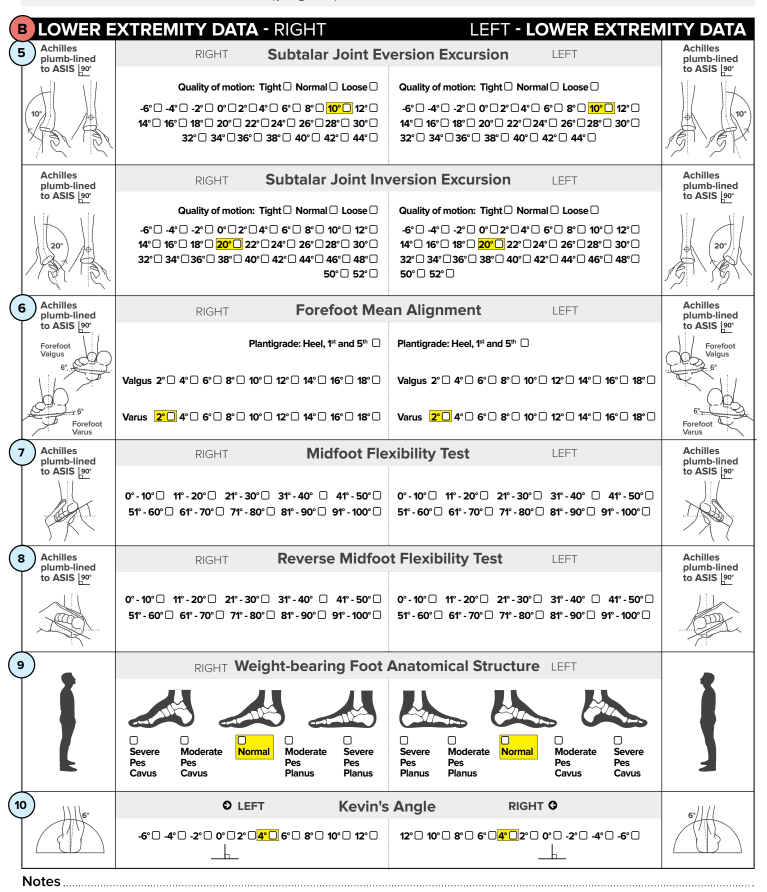
Biomechanical Evaluation (page 1)

A PATIENT			
First Name		Duration:	
Last Name		Onset:	
Sex: ☐ M ☐ F	Height Weight	Course	
Date of birth	//Subjective shoe size	Aggravating/Alleviating Factors:	
Diagnosis			
B LOWER E	XTREMITY DATA - RIGHT	LEFT - LOWER EXTREM	ITY DATA
1) _{[xi}	ASIS Wi	dth (cm)	ixi
	14	25 26 27 28 29 30 31 32 33 34 35 Average female	
Perform in Fowler or supine position	RIGHT External Hi	p Excursion LEFT	Perform in Fowler or supine position
Achilles Plumb-lined	Quality of motion: Tight ☐ Normal ☐ Loose ☐	Quality of motion: Tight ☐ Normal ☐ Loose ☐	Achilles Plumb-lined
Plumb-lined to ASIS 190°	0° 0 2° 0 4° 0 6° 0 8° 0 10° 0 12° 0 14° 0 16° 0 18° 0	0° \(2^\circ\) 4° \(\) 6° \(\) 8° \(\) 10° \(\) 12° \(\) 14° \(\) 16° \(\) 18° \(\)	Plumb-lined to ASIS [90]
	20° \(\) 22° \(\) 24° \(\) 26° \(\) 28° \(\) 30° \(\) 32° \(\) 34° \(\) 36° \(\) 38° \(\) 40° \(\) 42° \(\) 44° \(\) 46° \(\) 48° \(\) 50° \(\) 52° \(\)	20° \(22° \(\text{ 24° \(\text{ 26° \(\text{ 28° \(\text{ 30° \(\text{ 33° \(\text{ 34° \(\text{ 36° \(\text{ 38° \(\text{ 40° \(\text{ 44° \(\text{ 48° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 38° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 38° \(\text{ 36° \) 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36	
	54° 56° 58° 60° 62° 64° 66° 68° 70°	54° \[56° \] 58° \[60° \] 62° \[64° \] 66° \[68° \] 70° \[68° \]	
	72° 74° 76° 78° 80° 82° 84° 86° 88°	72° 🗆 74° 🗆 76° 🗆 78° 🗆 80° 🗆 82° 🗆 84° 🗆 86° 🗆 88° 🗆	
2 nd met	90° 92° 94° 96° 98° 100° 102° 104°	90° 92° 94° 96° 98° 100° 102° 104°	2 nd met
Vertical 2 nd met is 0° Perform in			Vertical 2 nd met is 0° Perform in Fowler or
Fowler or supine position	RIGHT Internal Hip Excursion LEFT		
- Starte	Quality of motion: Tight ☐ Normal ☐ Loose ☐	Quality of motion: Tight ☐ Normal ☐ Loose ☐	Achilles
Achilles Plumb- lined to	0° 0 2° 0 4° 0 6° 0 8° 0 10° 0 12° 0 14° 0 16° 0 18° 0	0° \(\) 2° \(\) 4° \(\) 6° \(\) 8° \(\) 10° \(\) 12° \(\) 14° \(\) 16° \(\) 18° \(\)	lined to
ASIS 90°	20° \[22° \] 24° \[26° \] 28° \[30° \] 32° \[34° \] 36° \[38° \] 40° \[42° \] 44° \[46° \] 48° \[50° \] 52° \[52° \]	20° \(22° \(\text{ 24° \(\text{ 26° \(\text{ 28° \(\text{ 30° \(\text{ 32° \(\text{ 34° \(\text{ 36° \(\text{ 38° \(\text{ 40° \(\text{ 42° \(\text{ 44° \(\text{ 48° \(\text{ 30° \(\text{ 50° \(\text{ 52° \(\text{ 36° \(\text{ 38° \(\text{ 30° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 38° \(\text{ 30° \(\text{ 36° \(\text{ 36° \(\text{ 36° \(\text{ 38° \(\text{ 30° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36° \(\text{ 36° \) 36	
	54° 56° 58° 60° 62° 64° 66° 68° 70°	54° \ 56° \ 58° \ 60° \ 62° \ 64° \ 66° \ 68° \ 70° \	
Jest Committee of the C	72° 🗆 74° 🗆 76° 🗆 78° 🗆 80° 🗆 82° 🗆 84° 🗆 86° 🗆 88° 🗆	72° 🗆 74° 🗆 76° 🗆 78° 🗆 80° 🗆 82° 🗆 84° 🗆 86° 🗆 88° 🗆	
÷ 2 nd met	90° 🗆 92° 🗆 94° 🗆 96° 🗆 98° 🗆 100° 🗆 102° 🗆	90° □ 92° □ 94° □ 96° □ 98° □ 100° □ 102° □	2 nd met
Vertical 2 nd met is 0° Fowlers, supine			Vertical 2 nd met is 0° Fowlers, supine or
or standing	RIGHT Genu \	Valgum LEFT	standing
Bisect Femur 8° 14°	-8° -6° -4° -2° 0° 2° 4° 6° 8° 1	-8° 🗆 -6° 🗀 -4° 🗀 -2° 🗀 0° 🗀 2° 🗀 4° 🗀 6° 🗆 8° 🗋	Bisect Femur
Bisect	10° □ 12° □ 14° □ 16° □ 18° □ 20° □ 22° □ 24° □ 26° □	10° □ 12° □ 14° □ 16° □ 18° □ 20° □ 22° □ 24° □ 26° □	Bisect
Tibia			Tibia
4	RIGHT Foot Dorsiflex	tion Excursion LEFT	Sec. Sec.
5 th met	Tot Dollar		5 th met
Tibia Silfverskiold Push Up			
	0° \(2° \) 4° \(\) 6° \(\) 8° \(\) 10° \(\) 12° \(\) 14° \(\) 16° \(\) 18° \(\)	0° \(2° \) 4° \(\) 6° \(\) 8° \(\) 10° \(\) 12° \(\) 14° \(\) 16° \(\) 18° \(\)	
	20° □ 22° □ 24° □ 26° □ 28° □ 30° □	20° □ 22° □ 24° □ 26° □ 28° □ 30° □	
5 th met			5 th met
Tibia 0°	Inverted Silfv	rerskiold Test -12° □ -10° □ -8° □ -6° □ -4° □ <mark>-2° □ 0° □</mark> 2° □ 4° □ 6° □	Tibia
- Jo	8° 10° 12° 14° 16° 18° 20°		
			





Biomechanical Evaluation (page 2)







Tests and Gait Evaluation

C	LOWER E	XTREMITY TESTING - RIGHT LEFT - LOWER EXTREMIT	TY TESTING		
11		RIGHT Extrinsic Muscle Testing LEFT			
	→	Invertors 5/5 4/5 3/5 2/5 1/5 0/5 0/5 2/5 2/5 3/5 4/5 5/5 0	\(\frac{1}{2}\)		
	ρη.) \	Evertors			
	NEW J	5/5 4/5 3/5 2/5 1/5 0/5 0/5 1/5 2/5 3/5 4/5 5/5 0			
	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	Dorsiflexors 5/5 4/5 3/5 2/5 1/5 0/5 0/5 1/5 2/5 3/5 4/5 5/5			
	↑				
	/ \ Plantar Flexors				
	₹	<u>5/5 4/5 3/5 2/5 1/5 0/5 </u> <u>0/5 1/5 2/5 3/5 4/5 5/5 </u>	+		
12		RIGHT Neurological Testing LEFT			
		Romberg			
	¥.	Balance intact ☐ Positive Romberg ☐ Positive Romberg ☐ Balance intact ☐	Ĩ		
		Patellar Reflex Absent despite reinforcement Absent despite reinforcement Present only with reinforcement Present only with reinforcement Normal Increased but normal Increased but normal Markedly hyperactive, with clonus Achilles Reflex Absent despite reinforcement Absent despite reinforcement Present only with reinforcement Present only with reinforcement Increased but normal Increased but normal Increased but normal Increased but normal Markedly hyperactive, with clonus Markedly hyperactive,			
	Positive Absent	Babinski Sign Absent □ Positive □ Positive □ Absent □	Absent Positive		
	Absent	Ankle Clonus Absent □ Positive (5 beats or more) □ Positive (5 beats or more) □ Absent □	Absent		
		Protective Sensation (Semmes Weinstein)	<i>Ŋ</i> _₩		
	17	Present ☐ Absent ☐ Present ☐ Present ☐			
C		Gait Evaluation			
		Gait Pattern Normal Hemiplegic Spastic Diplegic Neuropathic Myopathic Parkinsonian Choreiform Ataxic (cerebellar) Sensory Comment on head, shoulders, spine, pelvis, sagittal/transverse/frontal posture, etc.:			





Document of Medical Necessity Custom Molded Gauntlet

Patient Name:
HICN:
Prognosis: good
Duration of usage: 12 months
I certify that patient
Sufficient documentation of the medical condition found in the patient's medical record confirms the necessity for the type and quantity of items ordered. Therapy Goals (mark all that apply): To improve mobility To improve stability of lower extremity Decrease pain Aid healing soft tissue
☐ Promote immobilization, healing and treatment of injury
☑ NECESSITY OF ANKLE FOOT ORTHOSIS MOLDED TO PATIENT MODEL
The following criteria, which are specific to the condition of this patient, has justified the prescription of a custom ankle foot orthosis over prefabricated items. (mark all that apply): Patient could not successfully be fitted with a prefabricated AFO Condition necessitating the orthosis is expected to be permanent or lasting for a duration of more than 6 months Ankle or foot control is required on more than one plane of motion Adequate documentation of a neurological, circulatory, or orthopedic condition that requires custom fabrication has been provided by the patient Patient has display a lack of normal anatomical integrity or anthropometric proportions regarding a healing fracture
I hereby certify that an ankle foot orthotic described above is a semi-rigid or rigid device whose use is solely intended for the support of a weak or deformed limb or the restriction and/or elimination of motion in an injured and/or diseased part of the body. The AFO is designed to provide the appropriate stabilization, support and counterforces necessary for the limb or other extremity that requires bracing. All of the observations, physical exams, and documentation has provided me with enough sound evidence that it is my opinion that a custom molded ankle foot orthosis is both reasonable and necessary in reference to the accepted guidelines of medical practice in the treatment of the patient condition and rehabilitation.
Prescribing Clinician (printed name): Signature of Prescribing Clinician: Phone: Type I NPI:
Date:/





Prescription (Rx) Custom Molded Gauntlet

₱ BASIC INFORMATION:					
Clinician Name:		Patient Name:			
Duration of usag	ge:	Product Brand & Model:			
	T INCORMATION (MARK AND MEGES)				
	T INFORMATION (MARK ALL NECESSA				
□ G50 - Cabbie R □ L □ L1907	7 Ankle orthosis, supramalleolar with straps, with or	☐ G250 - Overlapping Articulating AFO R ☐ L ☐ L1970 Ankle foot orthosis, plastic with ankle joint, custom			
R□ L□ L2330	ut interface/pads, custom fabricated O Addition to lower extremity, lacer molded	fabricated R□L□ L2275 Addition to lower extremity, varus/valgus correction,			
to pat G100 - Short	tient model, for custom fabricated orthosis only AFO	plastic modification, padded/lined R□L□ L2330 Addition to lower extremity, lacer molded to patient			
	Ankle foot orthosis, plastic or other material, custom	model, for custom fabricated orthosis only R□L□ L2820 Addition to lower extremity, soft interface for molded			
R□ L□ L282	O Addition to lower extremity, soft interface for molded	plastic, below knee section			
R□ L□ L2330	c, below knee section O Addition to lower extremity, lacer molded	□ G300 - Tamarack Dorsi Assist AFO R □ L □ L1970 Ankle foot orthosis, plastic with ankle joint, custom			
to pat	tient model, for custom fabricated orthosis only	fabricated R □ L □ L2820 Addition to lower extremity, soft interface for molded			
R□ L□ L1940) Ankle foot orthosis, plastic or other rial, custom fabricated	plastic, below knee section R□L□ L2330 Addition to lower extremity, lacer molded			
R □ L □ L282	O Addition to lower extremity, soft interface	to patient model, for custom fabricated orthosis only			
R□L□ L2330	olded plsatic, below knee section O Addition to lower extremity, lacer molded to patient	R □ L □ L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), each joint			
mode □ G140 - Tall A	el, for custom fabricated orthosis only	R □ L □ L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), each joint			
R□ L□ L1940	Ankle foot orthosis, plastic or other material, custom	☐ G800 - Balance Brace AFO			
	O Addition to lower extremity, soft interface for molded	R □ L □ L1940 Ankle foot orthosis, plastic or other material, custom fabricated			
	c, below knee section O Addition to lower extremity, lacer molded to patient	R □ L □ L2820 Addition to lower extremity, soft interface for molded plastic, below knee section			
_ mode	el, for custom fabricated orthosis only rack Free Motion AFO	R □ L □ L2330 Addition to lower extremity, lacer molded to patient model, for custom fabricated orthosis only			
R□ L□ L1970	Ankle foot orthosis, plastic with ankle joint, custom	☐ Additional codes			
	O Addition to lower extremity, soft interface for molded	R □ L □ L2270 Addition to Lower Extremity Orthosis, Varus/Valgus Correction - TStrap			
	c, below knee section O Addition to lower extremity, lacer molded to patient	R □ L □ L2350 Addition to Lower Extremity Orthosis, Prosthetic Type, Socket, Molded to Patient Model			
mode	el, for custom fabricated orthosis only	R 🗆 L 🗖 L5000 Partial foot, shoe insert with longitudinal arch, toe filler			
5 DV (2222)					
	ALL THAT APPLY) Corresponds to Biomecha				
Adult Acquired Flatfoot (PTTD)	Flat foot [pes planus] (acquired) Spontaneous rupture of other tendons, ankle and foc	□ R - M21.41 □ L - M21.42 ot □ R - M66.871 □ L - M66.872			
riatioot (PTTD)	Sportaneous rupture or other tendoris, ankle and rocDisorder of ligament, ankle	\Box R - M00.071 \Box L - M00.072 \Box R - M24.271 \Box L - M24.272			
	Disorder of ligament, foot	□ R - M24.274 □ L - M24.275			
Lateral Ankle	Other acquired deformities of foot Other specific joint derangements of ankle, not elsev	□ R - M21.6X1 □ L - M21.6X2 /here classified □ R - M24.871 □ L - M24.872			
Instability					
Amputation	Acquired absence of great toe	□ R - Z89.411 □ L - Z89.412			
	Acquired absence of other toe(s) Acquired absence of foot	□ R - Z89.421 □ L - Z89.422 □ R - Z89.431 □ L - Z89.432			
Foot Drop	Foot Drop, acquired	□ R - M21.371 □ L - M21.372			
	: Hemiplegia - affecting [right/left] dominant side : Hemiplegia - affecting [right/left] non-dominant side	□ R - I69.951 □ L - I69.952 □ R - I69.953 □ L - I69.954			
DJD of Ankle	Primary osteoarthritis, ankle and foot	□ R - M19.071 □ L - M19.072			
and Rearfoot	Pain in ankle and joints of foot	□ R - M25.571 □ L - M25.572			
	Pain in lower leg Pain in foot	□ R - M79.661 □ L - M79.662 □ R - M79.671 □ L - M79.672			
	Other specified congenital deformities of feet	□ Q66.89			
	Other				
M THEDADY COAL(S): (MARK ALL THAT ARRIV)					
☑ THERAPY GOAL(S): (MARK ALL THAT APPLY) ☐ To improve mobility ☐ Promote healing soft tissue ☐ Decrease pain ☐ To improve stability of lower extremity ☐ Treatment of injury					
☐ To improve mobility ☐ Promote healing soft tissue ☐ Decrease pain ☐ To improve stability of lower extremity ☐ Treatment of injury					
•	9	Order/			
rype i NPI:		Date: MM DD YYYY			

1 CLINIC	CIAN REQUIRED		Kevin R	Root	KEVII ORTHOPED	N °
	Location		Tel: 1-800-496-0987 Fax: 1-866-919-9268		Gauntlet AFO	Order Form
			www.kevinrootmedica hello@kevinrootmedica		ORDER FORM F	PAGE 1 OF 1
Clinician Em	nail	<u></u>	Contact me to revi	ew Date:	//	□ Rush order due date: // MM DD YYYY
2 PATIEN	NT REQUIRED		3 SIDE	IMPR	SSION	FITTING
Patient's Em	ail		Use separate Rx for each side		Plaster	Shoe size:
First Name			Right	5 0	STS Casting Socks	Fit AFO to submitted:
Last Name					3D Foot Scanner	☐ Tracing
DOB/	/	Height	*G800 exempt BIL standard		Fiberglass Casting Tap	oe Shoes
4 AFO R	EQUIRED					
	Vacuum formed	Leather liner Lace closures Device to mets		• 4mm po	intrinsic post •	tion Gauntlet Leather liner Velcro closures Device to mets
	L Codes: L1907, L2330				L1970, L2330, L2820	Lutino Compilato
	4mm poly frame	Leather liner Optional closures Device to mets		9" heigh4mm po	lypropylene frame intrinsic post	lating Gauntlet 2 anterior/1 posterior straps Exterior struts Device to mets
0.00	L Codes: L1940, L2820, L2330			_	L1970, L2275, L2820, L23	
	4mm polypropylene frame	Leather liner Optional closures Device to mets		9" heigh4mm poRearfoot	lypropylene frame •	Leather liner Optional closures Device to mets
	L Codes: L1940, L2820, L2330				L1970, L2330, L2820, L22	
	 4mm polypropylene frame 	Leather liner Optional closures Device to mets		Send sh9" heigh4mm po	t .	t Gauntlet Leather liner Velcro and lace Fit to Shoes
	L Codes: L1940, L2820, L2330 G160 Only Leather Gaur	atlet		_	L1960, L5000, L2330, L2 Blance Brace Gau	
	9" heightNo frameNo post	Leather liner Optional closures Device to mets Custom scaphoid		Bilateral Si 9" heigh 3mm po Leather	tandard O Right Onl t lypropylene frame top cover	
5 IMPRE	SSION PREPARATION	PEOLIPED	A CLOSURE		L1940, L2820, L2330	S: OPTIONAL
Ankle	☐ As is ☐ Correct to ☐ 90°	×?	Lace/ velcro	Lace	Velcro	
Rearfoot	As is Correct to 90°		Color: Black	Brown		Taupe White
Forefoot	☐ As is ☐ Balance FF to RF	10 ×?	Full Plastazote I under leather	Linor –		OPTIONAL 6mm 9mm
	Special Instructions:		-			





Proof of Delivery - original in patient's chart, copy given to patient

SUPPLIER INFORMATION		
Supplier Name:		
HICN:		
THON		
PRODUCT INFORMATION (I	MARK ALL NECESSARY CODE	S AND ADDITIONS THAT APPLY)
G50 - Cabbie AFO		- Overlapping Articulating AFO
R 🗆 L 🗆 L1907 Ankle orthosis, supramalleola	r with straps, with or R 🗆 L 🗆	L1970 Ankle foot orthosis, plastic with ankle joint, custom
without interface/pads, custom fabric R L L L2330 Addition to lower extremity, la		fabricated L2275 Addition to lower extremity, varus/valgus correction,
to patient model, for custom fabricat	ted orthosis only	plastic modification, padded/lined
G100 - Short AFO	R D L D	L2330 Addition to lower extremity, lacer molded to patient
R □ L □ L1940 Ankle foot orthosis, plastic or fabricated		model, for custom fabricated orthosis only L2820 Addition to lower extremity, soft interface for molded
R□ L□ L2820 Addition to lower extremity, s	soft interface for molded	plastic, below knee section
plastic, below knee section R L L L2330 Addition to lower extremity, la	☐ G300 acer molded P☐ I ☐	- Tamarack Dorsi Assist AFO L1970 Ankle foot orthosis, plastic with ankle joint, custom
to patient model, for custom fabricat	ted orthosis only	fabricated
G120 - Standard AFO		L2820 Addition to lower extremity, soft interface for molded
R L L L1940 Ankle foot orthosis, plastic or material, custom fabricated	R [] I [plastic, below knee section L2330 Addition to lower extremity, lacer molded
R □ L □ L2820 Addition to lower extremity, s	soft interface	to patient model, for custom fabricated orthosis only
for molded plsatic, below knee secti R \subseteq L \subseteq L2330 Addition to lower extremity, la	on R∪L∪ acer molded to patient	L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), each joint
model, for custom fabricated orthosi		L2210 Addition to lower extremity, dorsiflexion assist (plantar
☐ G140 - Tall AFO R☐ L☐ L1940 Ankle foot orthosis, plastic or	other meterial quetem	flexion resist), each joint
fabricated	R L	 Balance Brace AFO L1940 Ankle foot orthosis, plastic or other material, custom
R□ L□ L2820 Addition to lower extremity, s	soft interface for molded	fabricated
plastic, below knee section R L L2330 Addition to lower extremity, la	R □ L □ acer molded to patient	L2820 Addition to lower extremity, soft interface for molded plastic, below knee section
_ model, for custom fabricated orthosi		L2330 Addition to lower extremity, lacer molded to patient
G200 - Tamarack Free Motion AFO	th ankle joint sustem	model, for custom fabricated orthosis only onal codes
R□L□ L1970 Ankle foot orthosis, plastic wit fabricated		L2270 Addition to Lower Extremity Orthosis, Varus/Valgus
R □ L □ L2820 Addition to lower extremity, s	soft interface for molded	Correction - TStrap
plastic, below knee section R L L2330 Addition to lower extremity, la	R □ L □ acer molded to patient	L2350 Addition to Lower Extremity Orthosis, Prosthetic Type, Socket, Molded to Patient Model
model, for custom fabricated orthosi	is only R□ L□	L5000 Partial foot, shoe insert with longitudinal arch, toe filler
sage Instructions:	to assist in immobilizing your foot and	ankle. Any AEO requires a period of adjustment (usually two
		Inkle. Any AFO requires a period of adjustment (usually two in the previous day. Please continue this process for two weeks.
		ning of the brace should occur, you may be walking too frequent
		ightness or uncomfortability continues, please contact the Kevin
thopedic office immediately. If the device cracks of	or breaks, remove it promptly and do no	ot wear again until you contacted our office immediately. Do not
	brace should be kept clean to ensure t	ne device can be properly secured. Skin moisturizes and knee
h socks can be used to prevent skin irritation.		
aterial failure warranty coverage:		
	vered at no-charge for up to 2 years.	all soft materials are covered at no-charge for up to ninety days.
cluding: material covers, Velcro straps, laces, and	l limb support pads.	
e item(s) that have been indicated. The supplier hap proper care for this item(s) will result in a voided w	as reviewed and provided me with writt varranty. A voided warranty as a result o	Medicare Supplier Standards. I certify that I have received en instructions of proper usage and care. I understand that of improper care could result in my responsibility for future r has instructed me to call the office if any difficulties or
with the device disc.		
Patient Name (printed)	Pa	tient Signature:
i atient maine (printed)		dent Signature
		Delivery//

Provide a copy to patient

Disclaimer: The codes within these pages are the offered suggestion based upon the HCPCS and ICD-10 codes provided by hcpcscodes.org and icdlist.com. Each prescribing practitioner should contact their local consultant or Medicare office to verify all billing codes, regulations and guidelines according to their geographic region.





MEDICARE DMEPOS SUPPLIER STANDARDS

Note: This is an abbreviated version of the supplier standards every Medicare DMEPOS supplier must meet in order to obtain and retain their billing privileges. These standards, in their entirety, are listed in 42 C.F.R. 424.57(c).

- 1. A supplier must be in compliance with all applicable Federal and State licensure and regulatory requirements.
- 2. A supplier must provide complete and accurate information on the DMEPOS supplier application. Any changes to this information must be reported to the National Supplier Clearinghouse within 30 days.
- 3. A supplier must have an authorized individual (whose signature is binding) sign the enrollment application for billing privileges.
- 4. A supplier must fill orders from its own inventory, or contract with other companies for the purchase of items necessary to fill orders.

A supplier may not contract with any entity that is currently excluded from the Medicare program, any State health care programs, or any other Federal procurement or non-procurement programs.

- 5. A supplier must advise beneficiaries that they may rent or purchase inexpensive or routinely purchased durable medical equipment, and of the purchase option for capped rental equipment.
- 6. A supplier must notify beneficiaries of warranty coverage and honor all warranties under applicable State law, and repair or replace free of charge Medicare covered items that are under warranty.
- 7. A supplier must maintain a physical facility on an appropriate site and must maintain a visible sign with posted hours of operation. The location must be accessible to the public and staffed during posted hours of business. The location must be at least 200 square feet and contain space for storing records.
- 8. A supplier must permit CMS or its agents to conduct on-site inspections to ascertain the supplier's compliance with these standards.
- 9. A supplier must maintain a primary business telephone listed under the name of the business in a local directory or a toll free number available through directory assistance. The exclusive use of a beeper, answering machine, answering service or cell phone during posted business hours is prohibited.
- 10. A supplier must have comprehensive liability insurance in the amount of at least \$300,000 that covers both the supplier's place of business and all customers and employees of the supplier. If the supplier manufactures its own items, this insurance must also cover product liability and completed operations.
- 11. A supplier is prohibited from direct solicitation to Medicare beneficiaries. For complete details on this prohibition see 42 CFR \S 424.57 (c) (11).
- 12. A supplier is responsible for delivery of and must instruct beneficiaries on the use of Medicare covered items, and maintain proof of delivery and beneficiary instruction.
- 13. A supplier must answer questions and respond to complaints of beneficiaries, and maintain documentation of such contacts.
- 14. A supplier must maintain and replace at no charge or repair cost either directly, or through a service contract with another company, any Medicare-covered items it has rented to beneficiaries.

- 15. A supplier must accept returns of substandard (less than full quality for the particular item) or unsuitable items (inappropriate for the beneficiary at the time it was fitted and rented or sold) from beneficiaries.
- 16. A supplier must disclose these standards to each beneficiary it supplies a Medicare-covered item.
- 17. A supplier must disclose any person having ownership, financial, or control interest in the supplier.
- 18. A supplier must not convey or reassign a supplier number; i.e., the supplier may not sell or allow another entity to use its Medicare billing number.
- 19. A supplier must have a complaint resolution protocol established to address beneficiary complaints that relate to these standards. A record of these complaints must be maintained at the physical facility.
- 20. Complaint records must include: the name, address, telephone number and health insurance claim number of the beneficiary, a summary of the complaint, and any actions taken to resolve it.
- 21. A supplier must agree to furnish CMS any information required by the Medicare statute and regulations.
- 22. All suppliers must be accredited by a CMS-approved accreditation organization in order to receive and retain a supplier billing number. The accreditation must indicate the specific products and services, for which the supplier is accredited in order for the supplier to receive payment for those specific products and services (except for certain exempt pharmaceuticals).
- 23. All suppliers must notify their accreditation organization when a new DMEPOS location is opened.
- 24. All supplier locations, whether owned or subcontracted, must meet the DMEPOS quality standards and be separately accredited in order to bill Medicare.
- 25. All suppliers must disclose upon enrollment all products and services, including the addition of new product lines for which they are seeking accreditation.
- 26. A supplier must meet the surety bond requirements specified in 42 CFR § 424.57 (d).
- 27. A supplier must obtain oxygen from a state-licensed oxygen supplier.
- 28. A supplier must maintain ordering and referring documentation consistent with provisions found in 42 CFR § 424.516(f).
- 29. A supplier is prohibited from sharing a practice location with other Medicare providers and suppliers.
- 30. A supplier must remain open to the public for a minimum of 30 hours per week except physicians (as defined in section 1848(j) (3) of the Act) or physical and occupational therapists or a DMEPOS supplier working with custom made orthotics and prosthetics. DMEPOS suppliers have the option to disclose the following statement to satisfy the requirement outlined in Supplier Standard 16 in lieu of providing a copy of the standards to the beneficiary. The products and/or services provided to you by (Kevin Orthopedic) are subject to the supplier standards contained in the Federal regulations shown at 42 Code of Federal Regulations Section 424.57(c). These standards concern business professional and operational matters (e.g. honoring warranties and hours of operation). The full text of these standards can be obtained at http://www.ecfr.gov. Upon request we will furnish you a written

copy of the standards.





Dispensing Documentation Custom Molded Gauntlet

₱ PATIENT INFORMATION	
Patient Name:	
HICN:	
PRODUCT INFORMATION (MARK ALL NECESSAI	RY CODES AND ADDITIONS THAT APPLY)
☐ G50 - Cabbie AFO	☐ G250 - Overlapping Articulating AFO
R 🗆 L 🗆 L1907 Ankle orthosis, supramalleolar with straps, with or	R 🗆 L 🗅 L1970 Ankle foot orthosis, plastic with ankle joint, custom
without interface/pads, custom fabricated R□ L□ L2330 Addition to lower extremity, lacer molded	fabricated R□ L□ L2275 Addition to lower extremity, varus/valgus correction,
to patient model, for custom fabricated orthosis only G100 - Short AFO	plastic modification, padded/lined R □ L □ L2330 Addition to lower extremity, lacer molded to patient
R 🗆 L 🗅 L1940 Ankle foot orthosis, plastic or other material, custom	model, for custom fabricated orthosis only
fabricated R□L□ L2820 Addition to lower extremity, soft interface for molded	R □ L □ L2820 Addition to lower extremity, soft interface for molded plastic, below knee section
plastic, below knee section	G300 - Tamarack Dorsi Assist AFO
R □ L □ L2330 Addition to lower extremity, lacer molded to patient model, for custom fabricated orthosis only	R □ L □ L1970 Ankle foot orthosis, plastic with ankle joint, custom fabricated
☐ G120 - Standard AFO	R □ L □ L2820 Addition to lower extremity, soft interface for molded
R 🗆 L 🗆 L1940 Ankle foot orthosis, plastic or other material, custom fabricated	plastic, below knee section $R \square L \square L2330$ Addition to lower extremity, lacer molded
R □ L □ L2820 Addition to lower extremity, soft interface for molded plsatic, below knee section	to patient model, for custom fabricated orthosis only R □ L □ L2210 Addition to lower extremity, dorsiflexion
R □ L □ L2330 Addition to lower extremity, lacer molded to patient	_ assist (plantar flexion resist), each joint
model, for custom fabricated orthosis only G140 - Tall AFO	R □ L □ L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), each joint
R 🗆 L 🗅 L1940 Ankle foot orthosis, plastic or other material, custom	G800 - Balance Brace AFO
fabricated R□L□ L2820 Addition to lower extremity, soft interface for molded	R □ L □ L1940 Ankle foot orthosis, plastic or other material, custom fabricated
plastic, below knee section	R □ L □ L2820 Addition to lower extremity, soft interface for molded
R □ L □ L2330 Addition to lower extremity, lacer molded to patient model, for custom fabricated orthosis only	plastic, below knee section $R \square L \square L2330$ Addition to lower extremity, lacer molded to patient
☐ G200 - Tamarack Free Motion AFO R☐ L☐ L1970 Ankle foot orthosis, plastic with ankle joint, custom	model, for custom fabricated orthosis only Additional codes
fabricated	R L L 2270 Addition to Lower Extremity Orthosis, Varus/Valgus
R L L2820 Addition to lower extremity, soft interface for molded plastic, below knee section	Correction - TStrap $R \square L \square L2350$ Addition to Lower Extremity Orthosis, Prosthetic
R □ L □ L2330 Addition to lower extremity, lacer molded to patient	Type, Socket, Molded to Patient Model
model, for custom fabricated orthosis only	R 🗆 L 🗇 L5000 Partial foot, shoe insert with longitudinal arch, toe filler
Note:	
${\bf S}{\bf)}$ At the time of the patient's visit, a custom fabricated AFO was disper	nsed and fitted. Although the patient is ambulatory, their condition
and related symptoms deem that this medical device is necessary as p	part of their treatment. It is anticipated that this device will functionally
benefit the patient. This custom device is appropriate and utilized in an	attempt to avert surgery and because a prefabricated device is
inappropriate for the patient's condition.	
O) The device appears to fit well and comfortably on the patient during	g a gait analysis.
A) Regarding the good fit, the patient was able to wear properly and ar	mbulate without concern or distress. This device's function is to assis
motion and provide ankle joint stabilization.	
P) The patient had received satisfactory information on the goals and f	functions of the device. Demonstrations of proper application, wear.
and care for the device were shown to the patient. The patient was tole	
of shoes, including: running shoes, lace and velcro-fastening shoes/bo	
sandals. The device was suitable for the patient's condition and not sul	
reviewed all precautions. Written instructions, warranty information and	
All and any questions were answered.	a deepy of Billian early standards were provided to the patient
All and any questions were unswered.	
Additional Notes:	
Print Supplier Name:	Supplier Signature:
Dispensing Date:/	