These documents have been provided by

WorryFreeDME

Custom Molded Gauntlet Compliance Documentation Packet

WorryFree DME Compliance Documentation Packet

To be completed by physician:

Biomechanical Evaluation Form (Medical Record Information)

Documents medical necessity

Document of Medical Necessity

- □ Justifies qualification for use of AF0
- Details reason for prefabricated versus custom device
- □ Justifies level of fitting (off-the-shelf versus custom-fitted)
- $\hfill\square$ Justifies code(s) selected

Prescription

- $\hfill\square$ Description of the items
- Patient Name
- □ Physician's printed name
- Diagnosis
- □ Physician's signature (no stamps allowed)
- □ Date (no stamps allowed)
- $\hfill\square$ Indication if right and / or left limb affected

To be given to Patient:

Proof of Delivery

- Patient Printed Name
- □ Date of delivery
- □ Item Description
- □ Item Code(s)
- Patient Signature
- Patient Address

DMEPOS Supplier Standards

To be completed by Supplier / Physician:

Dispensing Chart Notes

- Type of orthosis
- Describes method of fitting
- $\hfill\square$ Documents patient satisfaction
- * Confirms delivery of Supplier Standards





PedAlign







These documents have been provided by WorryFreeDME

Created by:

The American College of FOOT & ANKLE ORTHOPEDICS

Biomechanical Evaluation Form



Patient Name:					
Chief Complaint:					
History of problem:					
Nature of discomfort/pain					
Location (anatomic)					
,					
Duration					
Onset					
Course					
Aggravating and/or alleviating factors					
Left Stance Evaluation	n: Right	Normative values:	Treatments and response		
Angle of gait:→					
Base of gait:→ Foot appearance					
Tibial influence		0°-2° varus or valgus			
Relaxed calcaneal stance pos		0°			
Neutral calcaneal stance pos	ition (NCSP)	0°			
Non-Weight Bearing Evaluation: Limb length:→		Equal			
Hip sagittal plane-		Lyou			
Knee extended		Flexion 120°/extension 20-30°			
Knee flexed		Flexion 45-60°/extension 20-30°			
Hip transverse plane-		45° each direction			
Knee extended Knee flexed		45 each direction			
Hip frontal plane		45° each direction			
Knee sagittal plane		Flexion 120°/extension 0-10°			
Knee recurvatum		Absent			
Ankle sagittal plane-		Description 10°/ploptorflowion 40,70°			
Knee extended Knee flexed		Dorsiflexion 10°/plantarflexion 40-70° Dorsiflexion 10°/plantarflexion 40-70°			
Subtalar joint-					
Inversion		20°			
Eversion		10°			
Subtalar joint axis location Midtarsal joint		0°			
1 st ray range of motion		Dorsal & plantar excursion 5mm			
1 st MTPJ range of motion		Dorsal 65° or >unloaded/20-40° loaded			
Lesser MTPJ's					
Other comments:					
Muscle testing (extrinsics):					
Invertors		5/5: normal strength			
Evertors		5/5: normal strength			
Dorsiflexors		5/5: normal strength			
Plantarflexors Abduction		5/5: normal strength 5/5: normal strength			
Adduction		5/5: normal strength			
Neurological testing:					
Romberg→		Balance intact			
Patellar reflex		2+ normal			
Achilles reflex Babinski		2+ normal No hallux extension			
Clonus		Absent			
Protective sensation		Present			
Gait Evaluation -					
Gait pattern	o poluio orgittal/				
Comment on head/shoulders, spin transverse/frontal plane, postural, e	etc.				
Footgear (size/width, wear pat	tern(s))→				
Existing orthoses/type→ Weight→					
Height→					
Biomechanical assessment:		· · · · · · · · · · · · · · · · · · ·			
Treatment plan:					
Enter assistant name		Enter date of exam			
Signature of assistant		Signature of physician			

Save in patient's chart

The Medicare Program Integrity Manual, states that "For any DMEPOS item to be covered by Medicare, the patients medical record contains sufficient documentation of the patients medical condition to substantiate the necessity for the type and quantity of the items ordered.

These documents have been provided by

WorryFreeDME

Document of Medical Necessity: Custom Molded Gauntlet

Patient Name:	HICN:	
Prognosis: □ Good Duration of usage: □ 12 Months to I	ong term	
I certify that Mr. / Ms	-	
an ankle foot orthosis used during ambulation based on meetin	g all of the following criteria. The pat	ient is:
□ Ambulatory, and		
\square Has weakness or deformity of the foot and ankle, and		
\square Requires stabilization for medical reasons, and		
\square Has the potential to benefit functionally		
The patient's medical record contains sufficient documentation for the type and quantity of the items ordered.	of the patients medical condition to	substantiate the necessity
The goal of this therapy: (indicate all that apply)		
Improve mobility		
Improve lower extremity stability		
Decrease pain		
Facilitate soft tissue healing		
\Box Facilitate immobilization, healing and treatment of an injury		
Necessity of Ankle Foot Orthotic molded to patient model:		
A custom (vs. prefabricated) ankle foot orthosis has been prescribed of this patient. (indicate all that apply)	I based on the following criteria which a	re specific to the condition
The patient could not be fit with a prefabricated AFO		
\Box The condition necessitating the orthosis is expected to be per	manent or of longstanding duration (mo	re than 6 months)
\Box There is need to control the ankle or foot in more than one pla	ane	
The patient has a documented neurological, circulatory, or ort a model to prevent tissue injury	hopedic condition that requires custom	fabrication over
\square The patient has a healing fracture that lacks normal anatomic	al integrity or anthropometric proportior	IS
I hereby certify that the ankle foot orthotic described above is a rigid or semi-rigid or restricting or eliminating motion in a diseased or injured part of the body. It is braced. In my opinion, the custom molded ankle foot orthosis is both reasonable a of the patient condition and rehabilitation.	designed to provide support and counterforce	on the limb or body part that is being
Signature of Prescribing Physician:	Type I NPI: D	ate://
Printed Name of Prescribing Physician	Phone:	





B Langer Biomechanics

The OHI Family of Brands

PedAlign







Rx: Custom Molded Gauntlet

Doctor Name: Patient Name: ____ Product Brand and Model: Prognosis: Good Duration of usage: 12 Months Product Information (Check brand and model, cirlcle base code and addition(s)): ☐ Arizona Brace[®] Standard, Tall, AZ Sporty[™], AZ Breeze[™], Arizona Balance Brace[™] Arizona Brace[®] - Extended, Unweighting R 1 **L1940** Plastic orthosis, custom molded from a model of the patient. R 1 L1960 Plastic orthosis, custom molded from a custom fabricated, includes casting and cast preparation. model of the patient, custom fabricated, includes R L L2330 Addition to lower extremity, lacer molded to patient model casting and cast preparation. R L L2820 Addition to lower extremity orthosis, soft interface for mold R 1 L2330 Addition to lower extremity, lacer molded to patient model plastic below knee section L2820 Addition to lower extremity orthosis, soft R 1 Arizona Brace[®] - Articulated interface for mold plastic below knee section R 1 L1970 A semi-rigid molded plastic orthosis to hold the foot in neutral position ☐ Arizona Mezzo[™] (dorsi-plantar flexion), controls foot position, custom molded from a model **L1907** Ankle orthosis, supramalleolar, with straps. of the patient, custom fabricated, includes casting and cast preparation. R 1 with or without pads, custom fabricated R L2330 Addition to lower extremity, lacer molded to patient model L R 1 L2330 Addition to lower extremity, lacer molded R L2820 Addition to lower extremity orthosis, soft interface for mold 1 to patient model plastic below knee section ☐ Arizona Mezzo[™] - Partial Foot If Dorsiflex assist, ADD: L1907 Ankle orthosis, supramalleolar, with straps, R L2210 Addition to lower extremity, dorsiflexion assist R L with or without pads, custom fabricated (plantar flexion resist), (two per brace) □ AZ Slim[™] R L2330 Addition to lower extremity, lacer molded to patient model R L L1904 AFO molded ankle gauntlet R L5000 Partial foot, shoe insert, with longitudinal L L2330 Addition to lower extremity, lacer molded to patient model R L arch, toe filler L2820 Addition to lower extremity orthosis, soft interface for plastic R L below knee section DX: (indicate all that apply) - Corresponds to Biomechanical Examination Form PTTD **DJD of Ankle and Rearfoot** Amputation Spontaneous rupture of other tendons, ankle and foot Acquired absence of great toe Primary osteoarthritis, ankle and foot right (M66.871) left (M19.072) right (Z89.411) left (Z89.412) right (M19.071) left (M66.872) Pain in ankle and joints of foot Disorder of ligament, ankle Acquired absence of other toe(s) right (M24.271) left (M24.272) right (Z89.421) left (Z89.422) right (M25.571) left (M25.572) Acquired absence of foot Disorder of ligament, foot Pain in lower leg right (M24.274) right (Z89.431) left (Z89.432) left (M24.275) right (M79.661) left (M79.662) Other acquired deformities of foot Pain in foot **Foot Drop** Ieft (M79.672) right (M21.6X1) Ieft (M21.6X2) right (M79.671) Foot Drop, acquired right (M21.371) left (M21.372) Other specified congenital deformities of feet (Q66.89) Foot Risk / Imbalance Hemiplegia Muscle weakness, generalized (M62.81) affecting right dominant side (I69.951) Other Ataxic gait (R26.0) affecting left dominant side (I69.952) Difficulty in walking (R26.2) affecting right non-dominant side (I69.953) affecting left non-dominant side (I69.954) Unsteadiness on feet (R26.81) Other abnormalities of gait and mobility (R26.89) Lateral Ankle Instability Other specific joint derangements of ankle, not Condition is bilaterial elsewhere classified right (M24.871) left (M24.872) Therapeutic Objective(s): (indicate all that apply) Improve mobility Improve lower extremity stability Decrease pain Facilitate soft tissue healing Facilitate immobilization, healing and treatment of an injury

Signature of Prescribing Physician:			Type I NP	Type I NPI: Order Date:/				
Prescribing Phys	sician Printed Name: _							
		Th	e OHI Family of Bra	nds				
		B Langer	PedAlign	ROOMY	safestep	The Orthotic		





Arizona AF0 (877) 780-8382 SafeStep (866) 712-7837

Gauntlet AFO Collection

Arizona Brace® - Articulated

Arizona Brace®

No.



	🗌 Standard	🗌 Tall			
	Color: Sand	Black	U White	Brown	🗌 Pink
	Closure: 🗌 Laces	Velcro	🗌 Speed L	.aces 🗌 Be	oot Hooks
	Hinge: 🗌 Tamara	ack	🗌 Tamara	ck Dorsi - Assi	st
	Arizona Bra	ce®			
	🗌 Unweighti	ng (Proxin	nal ht. 1" bo	elow fibular	head)
	Extended	(Proximal	ht. 1" bel	ow fibular h	iead)
-	Color: Sand	Black	U White	Brown	🗌 Pink
	Closure: 🗌 Laces	Velcro	🗌 Speed I	Laces 🗌 Bo	ot Hooks
	(5" above and				
	Color: Sand	🗌 Black	U White	Brown	🗌 Pink
	Closure: 🗌 Laces	U Velcro	Speed I	Laces 🗌 Bo	oot Hooks
		(Please	note: No F	Plastic Shel	I)

Standard (5" above ankle) Tall (9" above ankle)

Color: Sand Black White Brown Pink

Closure: Laces Velcro Speed Laces Boot Hooks

(5" above ankle)						
Color: Sand	Black	White	Brown	🗌 Pink		
Closure: 🗌 Laces	Velcro	Speed L	aces 🗌 Bo	ot Hooks		



Arizona Mezzo™

 Standard
 Partial Foot

 Color:
 Sand
 Black
 White
 Brown

 Closure:
 Laces



AZ Breeze™

 □ Standard
 □ Tall

 Color:
 □ Sand
 □ Black

 Closure:
 □ Laces
 □ Velcro
 □ Speed Laces
 □ Boot Hooks



□ Arizona Balance Brace[™] Color: □ Sand □ Black

Closure: Laces Velcro

Full foot & removable insole options not available on ABB
Bundle with Apex Balance Shoe (ABS)
Gender: ______ Size: _____ Width: _____

4

BasisTM Slip-On (Four-Way Stretchable Footwear/AFO Companion. Developed to Extend Home and Indoor AFO & Orthotic Wear Time For Up To Ten Hours Per Day. Sold Only in Pairs). Color: Black Gender: Male Female Size:_____

(**B**Langer

Ship to address: 4825 East Ingram St. Mesa, AZ 85205 Fax: 480.222.1599

Dispense Date: _____ Work Order #:

Additional Charge options:

- □ Foot plate to end of toes (Our standard trim length is proximal to met heads)
- 🗌 Removable, multi density insole

Patient Inform	ation:] Right Foot	🗆 Left Foot	□ Bilateral
Patient Name:				
Height:	Weight:	Shoe Size	: Gene	der: 🗆 M 🔲 F
Dx:		D.0.B: _		
Shipping and B	illing Informa	tion:		
Bill to my accour		count #		-
Practitioner:				
Email: PO#:	Provide email to recei	ve an email alert on	ce this order has beer	n shipped.
Facility Name:				
Phone:				
Fax:				
Ship to address:				
Bill to address:				
Manufacturing	and shippin	g:		
MFG: 3 Business	Days (\$75.0	0) 🗌 7 Bi	usiness Days	s (\$50.00)
Ship:	3 Day Air	🗌 2 Day Air	🗆 Overnigt	nt
Other:	·····			
Special Instruc the cast set to Leave cast er	our recomme xactly as is	endations, pl	ease choose:	-
Correct Foref	ool to Neutral	🗀 otner _		

Remarks:





The OHI Family of Brands









Proof of Delivery: Custom Molded Gauntlet

Supplier Name:					HICN:			
	Product Information (Check brand and model, cirlcle base code and addition(s)): □ Arizona Brace [®] Standard, Tall, AZ Sporty [™] , AZ Breeze [™] , Arizona Balance Brace [™]						Brace® - Extended, Unweighting	
F	२ २ २	L L L	 L1940 Plastic orthosis, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation. L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for mold plastic below knee section 			L	L1960 Plastic orthosis, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation. L2330 Addition to lower extremity, lacer molded to patient model	
	\riz	ona	Brace [®] - Articulated		R	L	L2820 Addition to lower extremity orthosis, soft interface for mold plastic below knee section	
F	3	L	L1970 A semi-rigid molded plastic orthosis to hold the foot in neutral position (dorsi-plantar flexion), controls foot position, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation.		Ari z R R	zona L	Mezzo [™] L1907 Ankle orthosis, supramalleolar, with straps, with or without pads, custom fabricated L2330 Addition to lower extremity, lacer	
	२ २	L	L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for mold plastic below knee section			_	molded to patient model Mezzo ^{™ -} Partial Foot L1907 Ankle orthosis, supramalleolar, with	
_	3	L	If Dorsiflex assist, ADD: L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), (two per brace)		R	L	straps, with or without pads, custom fabricated L2330 Addition to lower extremity, lacer molded to patient model	
-	7	Slim' L	L1904 AFO molded ankle gauntlet		R	L	L5000 Partial foot, shoe insert, with longitudinal arch, toe filler	
-	7 7	L	L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for plastic below knee section					

Instructions For Use:

You have been dispensed this custom molded ankle orthosis to immobilize your foot and ankle. An AFO often requires a period of adjustment. It is best to wear it for one hour more each day and to continue this for two weeks. It should only be removed as specifically instructed. If the brace feels too tight, you may be walking too much. Get off your feet, loosen any straps and elevate your foot until the tightness resolves. If your symptoms do not resolve, please contact our office immediately. Should the device crack or break, remove it and do not use it again until you contact our office. Straps, laces should be kept clean of clothing fabric to insure the device is properly secured to your extremity. Applying a skin moisturizer and wearing knee high socks will prevent your skin from irritation.

Material failure warrantee coverage:

- · Hardware, plastic and metal component are covered at no-charge for six months.
- · All soft materials: material covers, Velcro straps and limb support pads, are covered at no - charge up to ninety days.

1

I have read the posted Complaint Resolution Policy and have been provided with a copy of the Medicare Supplier Standards. I certify that I have received the item(s) indicated. The supplier has reviewed the instructions for proper use and care and provided me with written instructions. I understand that failure to properly care for this item(s) will result in the warranty being voided. This could result in my responsibility for future repair or replacement costs if my insurance policy will not cover such costs. The supplier has instructed me to call the office if I have any difficulties or problems with the device.

Patient Signature ____

Printed Patient Name

Date Delivered:	/	/	/

1

Patient Address

Original in patient's chart, copy to patient

The codes contained herein are not the official position or endorsement of any organization or company. They are offered as a suggestion based upon input from previous customers. Each prescribing practitioner should contact his or her local carrier or Medicare office to verify billing codes, regulations and guidelines relevant to their geographic location.

The OHI Family of Brands













Medicare Supplier Standards

- 1. A supplier must be in compliance with all applicable Federal and State licensure and regulatory requirements and cannot contract with an individual or entity to provide licensed services.
- 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. An authorized individual (one whose signature is binding) must sign the application for billing privileges.
- 4. A supplier must fill orders from its own inventory, or must contract with other companies for the purchase of items necessary to fill the order. A supplier may not contract with any entity that is currently excluded from the Medicare program, any State health care programs, or from 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.
- A supplier must maintain a physical facility on an appropriate site. This standard requires that the location is 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.
- A supplier must permit CMS, or its agents to conduct on-site inspections to ascertain the supplier's compliance with these standards. The supplier location must be accessible to beneficiaries during reasonable business hours, and must maintain a visible sign and posted hours of operation.
- 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 must agree not to initiate telephone contact with beneficiaries, with a few exceptions allowed. This standard prohibits suppliers from contacting a Medicare beneficiary based on a physician's oral order unless an exception applies.
- 12. A supplier is responsible for delivery and must instruct beneficiaries on use of Medicare covered items, and maintain proof of delivery.
- 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 directly, or through a service contract with another company, 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 supplier standards to each beneficiary to whom it supplies a Medicare covered item.
- 17. A supplier must disclose to the government any person having ownership, financial, or control interest in the supplier.
- 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 implementing 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 of those specific products and services (except for certain exempt pharmaceuticals). Implementation Date October 1, 2009
- 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.
- Must meet the surety bond requirements specified in 42 C.F.R. 424.57(c). Implementation date May 4, 2009
- 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 C.F.R. 424.516(f).
- 29. DMEPOS suppliers are prohibited from sharing a practice location with certain other Medicare providers and suppliers.
- 30. DMEPOS suppliers must remain open to the public for a minimum of 30 hours per week with certain exceptions.





The OHI Family of Brands

Blanger PedAlign







Dispensing Chart Notes: Custom Molded Gauntlet

Patient Name: HICN:							
			ormation (Check brand and model, cirlcle base code and addition(s)): Brace® Standard, Tall, AZ Sporty™, AZ Breeze™, Arizona Balance Brace™		Ari	zona	Brace® - Extended, Unweighting
	R R R	L L L	 L1940 Plastic orthosis, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation. L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for mold plastic below knee section 		R R	L	L1960 Plastic orthosis, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation. L2330 Addition to lower extremity, lacer molded to patient model
	Ariz	ona	Brace [®] - Articulated		R	L	L2820 Addition to lower extremity orthosis, soft
	R	L	L1970 A semi-rigid molded plastic orthosis to hold the foot in neutral position (dorsi-plantar flexion), controls foot position, custom molded from a model of the patient, custom fabricated, includes casting and cast preparation.		Ari R	zona L	interface for mold plastic below knee section Mezzo [™] L1907 Ankle orthosis, supramalleolar, with straps, with or without pads, custom fabricated
	R R	L	L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for mold plastic		R	L	L2330 Addition to lower extremity, lacer molded to patient model
	below knee section				Ari	Mezzo ^{™ -} Partial Foot	
	R	L	If Dorsiflex assist, ADD: L2210 Addition to lower extremity, dorsiflexion assist (plantar flexion resist), (two per brace)		R R	L	L1907 Ankle orthosis, supramalleolar, with straps, with or without pads, custom fabricated L2330 Addition to lower extremity, lacer
	AZ S	Slim'					molded to patient model
	R R R	L L L	 L1904 AFO molded ankle gauntlet L2330 Addition to lower extremity, lacer molded to patient model L2820 Addition to lower extremity orthosis, soft interface for plastic below knee section 		R	L	L5000 Partial foot, shoe insert, with longitudinal arch, toe filler

- S) A custom molded gauntlet was dispensed and fit at this visit. Patient is ambulatory. Due to the patient's condition and related symptoms this device is medically necessary as part of the overall treatment. It is anticipated that the patient will benefit functionally with the use of this device. The custom device is utilized in an attempt to avoid the need for surgery and because a prefabricated device is inappropriate.
- 0) Upon gait analysis, the device appeared to be fitting well and the patient states that the device is comfortable.
- A) Good fit. The patient was able to apply properly and ambulate without distress. The function of this device is to restrict and limit motion and provide stabilization in the ankle joint.
- P) The goals and function of this device were explained in detail to the patient. The patient was shown how to properly apply, wear, and care for the device. It was explained that the device will fit and function best in a lace-up shoe with a firm heel counter and a wide base of support. When the device was dispensed, it was suitable for the patient's condition and not substandard. No guarantees were given. Precautions were reviewed. Written instructions, warranty information and a copy of DMEPOS Supplier Standards were provided. All questions were answered.

Additional Notes: ________ Dispensing Date: _______ Dispensing Date: _______ Print Supplier Name: _______ The OHI Family of Brands _______

PedAlign











