Doctor Name: ___________________________________
Phone: ____________________________

Indicate Quantity: Bilateral

MBB
____ L1940 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.

____ L2330 Addition to lower extremity, lacer molded to patient model

____ L2820 Addition to lower extremity orthosis, soft interface for mold plastic below knee section

Dx: (check all that apply)

FALL RISK/IMBALANCE
❑ At Risk/History or Fall (V15.88)
❑ Muscle weakness (728.87)
❑ Ataxia, muscular incoordination (781.3)
❑ Gait abnormality/ staggering, ataxic (781.2)

DJD OF ANKLE AND REARFOOT
❑ Osteoarthritis, Localized Primary Ankle & Foot (715.17)
❑ Arthropathy, unspecified, ankle and foot (716.97)
❑ Pain in joint, ankle, foot (719.47)
❑ Ankle pain & support (729.5)

LATERNAL ANKLE INSTABILITY
❑ Instability of Joint, Ankle & Foot (718.87)

DROPFOOT
❑ Dropfoot (736.79)
❑ Hemiplegia (438.20)

THERAPEUTIC OBJECTIVE(S): (check all that apply)
❑ Improve mobility
❑ Improve lower extremity stability
❑ Decrease pain
❑ Reduce Risk of Falls
❑ Facilitate muscular coordination and gait stability
❑ Improve postural sway and ankle stability

Duration of usage: 12 Months

Signature of Prescribing Physician: ___________________________
Type I NPI: ___________________________ Date: _____/_____/_____

Custom MBB Compliance Documentation
**Document of Medical Necessity: Custom Molded Gauntlet Ankle Foot Orthotic**

**Patient Name:** __________________________________________

**HICN:** _______________ **DOB:** _____/_____/_____

**Prognosis:** Good  
**Duration of usage:** 12 Months

**Select Quantity, Product and HCPC Codes**

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1940</td>
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<td>Addition to lower extremity orthosis, soft interface for mold plastic below knee section</td>
</tr>
</tbody>
</table>

**Product Name:** __________________________________________

I hereby certify that Mr. / Ms. ____________________ ____________________ qualifies for and will benefit from the product designated above based on the following criteria (check all that apply):

- Partial or complete paralysis of one or more leg muscles.
- Significant weakness, ataxia or gait abnormality
- History of Falls
- Significant impairment of gait due to pain or ankle / foot deformity.
- Instability in gait with recurrent sprains or falls.

**The goal of this therapy:** (check all that apply)

- Improve mobility
- Improve lower extremity stability
- Decrease pain
- Decrease risk for fall

**Necessity of Ankle Foot Orthotic molded to patient model:**

A custom (vs. prefabricated) ankle foot orthosis has been prescribed based on the following criteria which are specific to the condition of this patient. (Check all that apply)

- The patient could not be fit with a prefabricated AFO
- The condition necessitating the orthosis is expected to be permanent or of longstanding duration (more than 6 months)
- There is need to control the ankle or foot in more than one plane
- The patient has a documented neurological, circulatory, or orthopedic condition that requires custom fabrication over a model to prevent tissue injury

I hereby certify that the ankle foot orthotic described above is a rigid or semi-rigid device which is used for the purpose of supporting a weak or deformed body member or restricting or eliminating motion in a diseased or injured part of the body. It is designed to provide support and counterforce on the limb or body part that is being braced. In my opinion, the custom molded ankle foot orthosis is both reasonable and necessary in reference to accepted standards of medical practice in the treatment of the patient condition and rehabilitation.

**Signature:** ________________________________  **Type I NPI:** _________  **Date:** _____/_____/_____

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**L2330** Addition to lower extremity, lacer molded to patient model

**L2820** Addition to lower extremity orthosis, soft interface for mold plastic below knee section
Dispensing Chart Notes: Custom Molded Gauntlet Ankle Foot Orthotic

Patient Information:
Mr. / Ms. ___________________________________________  HICN: __________________  DOB: ____/____/____

Dx: (check all that apply)

FALL RISK/IMBALANCE
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- Dropfoot (736.79)
- Hemiplegia (438.20)

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Quantity, Product and HCPC Codes

S) (Product name)___________________________________ was dispensed and fit at this visit. Patient is ambulatory. There is instability with range of motion that requires stabilization. Due to the indicated diagnosis(s) and related symptoms this device is medically necessary as part of the overall treatment. The function of this device is to stabilize gait, improve postural balance/sway, provide stabilization in the ankle joint and to reduce the risk for fall. It is anticipated that the patient will benefit functionally with the use of this device. The custom device is utilized in an attempt to reduce the mortality/morbidity of falling.

O) 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.

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 stiff 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 30 Supplier Standards were provided. All questions were answered.

Supplier Signature: ________________________________  Date: _____/_____/_____

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L1940  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.

L2330  Addition to lower extremity, lacer molded to patient model

L2820  Addition to lower extremity orthosis, soft interface for mold plastic below knee section
Patient Receipt: Custom Molded Ankle Foot Orthotic

Dr. _______________________ Phone: ______________________

Patient Information:
Mr./ Ms. ____________________ ____________________ HICN: ___________ DOB: __/__/____

Quantity, Product and HCPC Codes

___ L1940 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.
___ L2330 Addition to lower extremity, lacer molded to patient model
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Instructions For Use:

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.

You have been dispensed this custom molded ankle brace to stabilize your foot and ankle in order to prevents falls and imbalance. 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.

I have read the posted Complaint Resolution Policy and have been provided with a copy of the 30 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: _________________________ Date: _____/_____/______