



## The Equinus Brace™ Billing, Coding and Documentation Information Sheet

The Equinus Brace™ is treating equinus deformity and/or plantar fasciitis according to the LCD for the CPT code.

The LCD for L4396 consists of 5 requirements from a documentation standpoint.

A static ankle-foot orthosis (AFO) (L4396) is covered if **either all of criteria 1 – 4 or criterion 5 is met** and documented accordingly:

- 1) Plantar flexion contracture of the ankle with dorsiflexion on passive range of motion testing of at least 10 degrees (i.e., a non-fixed contracture); and,
- 2) Reasonable expectation of the ability to correct the contracture; and,
- 3) Contracture is interfering or expected to interfere significantly with the patient's functional abilities; and,
- 4) Used as a component of a therapy program which includes active stretching of the involved muscles and/or tendons.
- 5) The patient has plantar fasciitis

M24.571 – Contracture, Right Ankle - applies to criteria 1 - 4

M24.572 – Contracture, Left Ankle - applies to criteria 1 - 4

M72.2 - Plantar Fascia Fibromatosis (including Plantar Fasciitis) - applies to criteria 5

When the brace is being used for any pathology that has an associated equinus deformity (i.e. Achilles Tendonitis, Posterior Tibial Tendon Dysfunction, Metatarsalgia, Pronation Syndrome, Acquired Flatfoot Deformity), the diagnosis code for ankle contracture (i.e. M24.571/M24.572) should be linked to the CPT code for the brace.

If the brace is being used for plantar fasciitis without an equinus deformity, it should be linked to the plantar fasciitis diagnosis code (i.e. M72.2).

If the brace is being used for plantar fasciitis with an ankle contracture deformity, it can be linked with either diagnosis code or both codes (M24.571/M24.572/M72.2).

As always documentation is critical, especially for the ankle contracture deformity diagnosis (M24.571/M24.572). Please note that the ICD 10 for Short Achilles Tendon Acquired (M67.01/M67.02) are not listed in the LCD for L4396, therefore it is not recommended to utilize these codes with the CPT code for The Equinus Brace™. Here is a sample of suggested documentation.

### **Ankle Joint Objective Findings Documentation:**

Ankle joint dorsiflexion with the knee extended was measured with a goniometer/tractograph: right = \_\_\_\_ degrees and left = \_\_\_\_ degrees (abnormal considered less than 5 degrees)

Ankle joint dorsiflexion with the knee flexed was measured with a goniometer/tractograph: right = \_\_\_\_ degrees and left = \_\_\_\_ degrees (abnormal considered less than 10 degrees)

Total ankle joint range of motion measured with a goniometer/tractograph: Plantarflexion: right = \_\_\_\_ degrees and left = \_\_\_\_ degrees (40 – 60 degrees = normal); Dorsiflexion: right = \_\_\_\_ degrees and left = \_\_\_\_ degrees (10 – 20 degrees = normal)

There should also be documentation of symptoms and findings of with the associated lower extremity



pathological condition (i.e. Achilles tendonitis, Posterior Tibial Tendon Dysfunction, Metatarsalgia, Pronation Syndrome, Acquired Flatfoot Deformity) in the Musculoskeletal Objection Findings Documentation.

It is also important to document any prior treatment, especially manual stretching/physical therapy/shoe modifications in the History of Present Illness in the Subjective Complaint section of the note.

In the Plan section of the note, the suggested documentation for the brace is as follows.

### **The Equinus Brace™ Dispensing Documentation:**

A plastic prefabricated static Ankle Foot Orthosis was dispensed and fitted for the [right, left, bilateral] foot/feet at this visit. The device will be utilized for the next 8 to 12 weeks. Due to the pain in the foot/ankle/leg with weight bearing throughout the day, with diagnosis of equinus deformity and related symptoms, this is medically necessary for the treatment. The function of this device is to serve as an anticontracture device of the Gastrocsoleal complex and to restrict and limit motion and help reduce excessive stress and strain to Gastrocsoleal complex and foot/ankle/leg. It is being utilized to prevent the plantar contracture of the Gastrocsoleal complex. The goals of this therapy are to: 1. Treat plantarflexion contracture of the ankle with dorsiflexion, the ankle on passive range of motion testing is noted to have at least 10 degrees (i.e., a non-fixed contracture); and, 2. Provide a reasonable expectation of the ability to correct the contracture; and, 3. Reduce the contracture that is interfering or expected to interfere significantly with the beneficiary's functional abilities; and, 4. Used as a component of a therapy program which includes active stretching of the involved muscles and/or tendons; and/or 5. Treat the beneficiary's plantar fasciitis. Additionally, medial and lateral ankle dorsiflex assistive and plantarflex restraint hinges are included on the brace and were set at [ ] degrees of dorsiflexion. The intent of these hinges is to resist plantarflexion and assist with dorsiflexion of the Gastrocsoleal complex and/or plantar fascia. These hinges will be adjusted over the course of the patient's therapy. The patient was instructed to not adjust the hinges. They were also advised to bring the device to the office at their next visit for further evaluation and adjustments. The goals and function of this device were explained in detail to the patient. The patient states that the device is comfortable when applied. The patient was shown and told in detail how to properly wear and care for the device. The patient was able to apply and remove the device properly without assistance and to not ambulate with the device in place. The device was then dispensed and was suitable for the condition and not substandard. No guarantees regarding resolution of symptoms were given and precautions were reviewed. Written instructions and warranty information were given along with the list of the current Durable Medical Equipment Supplier Standards. The patient signed a written proof of delivery. All questions were answered to their satisfaction.

When billing for the The Equinus Brace™ the following is suggested.

L4396 - RT, LT (if using bilateral both RT, LT modifiers must be included).

However, it is important to note that it is best that they are billed separately i.e. L4396 RT billed as 1 unit and L4396 LT billed as 1 unit. This helps to insure accurate payment is made on both.

L2210 - RT, LT **EXCEPT for MEDICARE/GOVERNMENTAL PLANS** (The Equinus Brace is currently appealing this code with Medicare, this code may continue to be billed with non-governmental insurance provider), if using bilateral both RT, LT modifiers must be included AND the number of units per side (i.e. 2 per side as Equinus Brace™ has 2 hinges per brace). Again, it is important to note that it is best that each "pair" of hinges are billed separately i.e. L2210 RT billed as 2 units and L2210 LT billed as 2 units. This helps to insure accurate payment is made on all 4 hinges.

**\*\*\*Disclaimer\*\*\***

**The final and sole responsibility for the correct coding, within established laws, rules, and standards of practice, rests upon the party submitting the claim.**