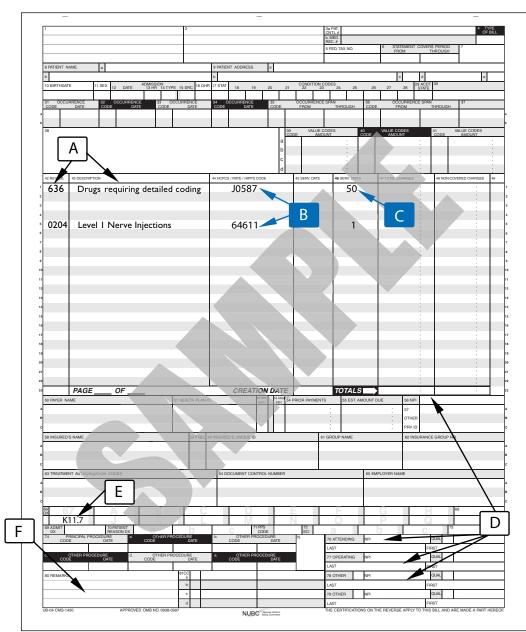
# SAMPLE UB-04 CLAIM FORM



## For Product Administered in the Hospital Outpatient Setting— Effective 8/19/19



Field 42 & 43: Enter the appropriate revenue codes & descriptions corresponding to HCPCS codes in Field 44 – eg: 0204 - Level 1 Nerve Injections 636 – Drugs requiring detailed coding 761 – Treatment Room Field 44: Enter the appropriate HCPCS and CPT codes: MYOBLOC – J0587, Botulinum Toxin Type B (per 100 Units) Injection – 64611, Chemodenervation of parotid and submandibular salivary glands, bilateral. Other administration codes may be appropriate Field 46:  $\sim$ Enter the number of billing Units. For J0587, a billing Unit is per 100 Units of MYOBLOC Please note that not all claims processing systems allow three digits in this field. In these cases, Units administered that are equal to or greater than 10,000 may need to be broken down on multiples lines (eg, 99, 98, and 3 for 20,000 Units). This billing example is for 5,000 Units. Fields 56, 76-79: D National Provider Identifier (NPI) Filed 56: Enter NPI for the Facility Field 76: Enter NPI for the Attending Physician Field 77: Enter NPI for the Operating Physician Field 78 and 79: Enter NPI for Other Provider Type Fields 67-75: Enter the ICD-10-CM (10) diagnosis code that is appropriate for the patient. The

that is appropriate for the patient. The diagnosis code for disturbances of salivary secretion (sialorrhea) is K11.7. Other diagnosis codes may be acceptable. Please note that Field 67 is for the principal diagnosis and Fields 68-75 are for secondary diagnosis, if necessary.

### Field 80:

- Some payers may require that NDC numbers be entered into the electronic comment field. If required, the NDC numbers are entered with a "0" in the sixth position. See below:
- 10454-0710-10 MYOBLOC 2,500 Units/0.5 mL
- 10454-0711-10 MYOBLOC 5,000 Units/1 mL
- 10454-0712-10 MYOBLOC 10,000 Units/2 mL

The above diagnosis and procedure codes are provided as examples only. The healthcare provider is responsible for determining the appropriate codes for an individual patient.

Please see Important Safety Information and Indication <u>(on Page 2)</u> and full <u>Prescribing Information</u>, including Boxed WARNING and <u>Medication Guide</u>.

### MYOBLOC is ready to use and requires no mixing

### 3 SINGLE-DOSE VIAL SIZES\*

\*For single-patient use. The recommended dosage of MYOBLOC for chronic sialorrhea is 1,500 to 3,500 Units; 500 to 1,500 Units per parotid gland and 250 Units per submandibular gland; no more frequent than every 12 weeks.



For more information on MYOBLOC Reimbursement Services, call 1-888-461-2255, Option 3, or visit <u>myobloc.com</u>

### **IMPORTANT SAFETY INFORMATION**

### WARNING: DISTANT SPREAD OF TOXIN EFFECT

See full prescribing information for complete boxed WARNING.

The effects of MYOBLOC® and all botulinum toxin products may spread from the area of injection to produce symptoms consistent with botulinum toxin effects. These symptoms have been reported hours to weeks after injection. Swallowing and breathing difficulties can be life threatening and there have been reports of death. The risk of symptoms is probably greatest in children treated for spasticity but symptoms can also occur in adults, particularly in those patients who have an underlying condition that would predispose them to these symptoms.

### CONTRAINDICATIONS

MYOBLOC is contraindicated in patients with:

- A known hypersensitivity to any botulinum toxin product or to any of the components in the formulation
- Infection at the proposed injection site(s)

#### WARNINGS AND PRECAUTIONS

Lack of Interchangeability Between Botulinum Toxin Products
 The potency units of MYOBLOC are specific to the preparation and
 biological activity assay method utilized. Due to differences in the aspects
 of this assay such as the vehicle, dilution scheme, and laboratory protocols
 for various potency assays, potency units are not interchangeable with
 other preparations of botulinum toxin products and, therefore, units of
 biological activity of MYOBLOC cannot be compared to or converted into
 units of any other botulinum toxin products assessed with any other

## specific assay method.Hypersensitivity Reactions

Serious hypersensitivity reactions have been reported with botulinum toxin products. Angioedema, urticaria, and rash have occurred with MYOBLOC treatment. Hypersensitivity reactions can also include anaphylaxis, serum sickness, soft tissue edema, and dyspnea. If serious and/or immediate hypersensitivity reactions occur, discontinue further injection of MYOBLOC and institute appropriate medical therapy immediately. The use of MYOBLOC in patients with a known hypersensitivity to any botulinum neurotoxin or to any of the excipients (human albumin, sucrose), could lead to a life-threatening allergic reaction.

### Dysphagia and Breathing Difficulties

Treatment with MYOBLOC and other botulinum toxin products can result in swallowing or breathing difficulties. Patients with pre-existing swallowing or breathing difficulties may be more susceptible to these complications. In most cases, this is a consequence of weakening of muscles in the area of injection that are involved in breathing or swallowing. When distant effects occur, additional respiratory muscles may be involved. Deaths as a complication of severe dysphagia have been reported after treatment with botulinum toxin. Dysphagia may persist for several months and require use of a feeding tube to maintain adequate nutrition and hydration. Aspiration may result from severe dysphagia and is a particular risk when treating patients in whom swallowing or respiratory function is already compromised.

### - Pre-Existing Neuromuscular Disorders:

Individuals with peripheral motor neuropathic diseases, amyotrophic lateral sclerosis, or neuromuscular junctional disorders (e.g., myasthenia gravis or Lambert-Eaton syndrome) should be monitored particularly closely when given botulinum toxin. Patients with neuromuscular disorders may be at increased risk of clinically significant effects including severe dysphagia and respiratory compromise from typical doses of MYOBLOC.

### Human Albumin and Transmission of Viral Diseases

This product contains albumin, a derivative of human blood. Based on effective donor screening and product manufacturing processes, it carries an extremely remote risk for transmission of viral diseases and variant Creutzfeldt-Jakob disease (vCJD). There is a theoretical risk for transmission of Creutzfeldt-Jakob disease (CJD), but if that risk actually exists, the risk of transmission would be considered extremely remote. No cases of transmission of viral diseases, CJD, or vCJD have ever been identified for licensed albumin or albumin contained in other licensed products.

### MOST COMMON ADVERSE REACTIONS (>5% of patients and >5% more than placebo)

Chronic Sialorrhea: dry mouth, dysphagia

### DRUG INTERACTIONS

### Aminoglycosides and Other Agents Interfering with Neuromuscular

Transmission: Co-administration of MYOBLOC and aminoglycosides or other agents interfering with neuromuscular transmission (e.g., curare-like compounds) should only be performed with caution as the effect of the toxin may be potentiated. Anticholinergic Drugs: Use of anticholinergic drugs after administration of MYOBLOC may potentiate systemic anticholinergic effects. Other Botulinum Neurotoxin Products: The effect of administering different botulinum toxin products at the same time or within several months of each other is unknown. Excessive neuromuscular weakness may be exacerbated by administration of another botulinum toxin prior to the resolution of the effects of a previously administered botulinum toxin. Muscle Relaxants: Excessive weakness may also be exaggerated by administration of a muscle relaxant before or after administration of MYOBLOC.

### INDICATION

MYOBLOC<sup>®</sup> injection is indicated for:

• the treatment of chronic sialorrhea in adults

### Please refer to the full <u>Prescribing Information</u> and <u>Medication Guide</u>.



