



Blue Cross  
Blue Shield  
Blue Care Network  
of Michigan

Nonprofit corporations and independent licensees  
of the Blue Cross and Blue Shield Association



**PAIN MANAGEMENT  
EPIDURAL STEROID INJECTIONS  
AUTHORIZATION REQUEST FORM**

Utilization management toll-free phone: 1-833-217-9670  
Utilization management local phone: 313-908-6040  
Utilization management fax: 313-483-7323

<b>Today's date (mm/dd/yyyy):</b> ___ / ___ / ____
<b>Provider contact name:</b>
<b>Provider contact phone:</b>
<b>Provider contact fax:</b>
<b>Provider contact email:</b>
<b>Provider name:</b>
<b>Provider TIN:</b>
<b>Provider NPI:</b>
<b>Practice/group name:</b>
<b>Provider physical address:</b>
<b>Provider mailing address (if different):</b>

<b>Member name:</b>
<b>Date of birth (mm/dd/yyyy):</b> ___ / ___ / ____
<b>Member ID (including any alpha prefix):</b>
<b>Health plan:</b>
<b>Notification method preference:</b> <input type="checkbox"/> Postal mail <input type="checkbox"/> Fax
<b>Mailing address or fax number:</b>
<b>Notes:</b>

<b>Where will the procedure take place?</b> <input type="checkbox"/> Provider office <input type="checkbox"/> Outpatient facility <input type="checkbox"/> Inpatient hospital <input type="checkbox"/> Ambulatory surgical center	
<b>Facility name:</b>	<b>Facility contact name:</b>
<b>Facility TIN:</b>	<b>Facility contact phone:</b>
<b>Facility NPI:</b>	<b>Facility contact fax:</b>
<b>Facility physical address:</b>	<b>Facility mailing address (if different):</b>



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Requested procedure code	Modifier: LT, RT or 50 (bilateral)	Quantity	Spine level

<b>Diagnosis code(s):</b>	<b>Anticipated date of service (mm/dd/yyyy):</b> ___ / ___ / ___
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**Case urgency**  
 Standard       Expedited

In keeping with guidelines from the National Committee for Quality Assurance and Centers for Medicare & Medicaid Services, prior authorization requests qualify for expedited review when the standard review time frame could do one of the following:

- Seriously jeopardize the life, health or safety of the member or others, due to the member's psychological state.
- In the opinion of a practitioner with knowledge of the member's medical or behavioral health condition, subject the member to adverse health consequences without the care or treatment that is the subject of the request.

<b>Patient's height:</b>	<b>Patient's weight:</b>	<b>Patient's BMI:</b>
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**What type of procedure is planned?** (Select one and answer all adjacent questions.)

<input type="checkbox"/> <b>Initial epidural steroid injection</b> (answer a – h)	a. Is there presence of moderate to severe pain (rated at least 3 out of 10) that interferes with daily activities?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	b. Are any radiculopathy or claudication symptoms (burning, tingling, cramping) present?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	c. Does imaging show stenosis or disc herniation/bulging that seems to match up with radiculopathy?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	d. Is the procedure planned for one caudal or interlaminar, or one bilateral or two unilateral transforaminal levels?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	e. Will more than the recommended amount of steroid be injected? (80mg of Triamcinolone, 80mg of methylprednisolone, 12mg of betamethasone, or 15mg of dexamethasone)	<input type="checkbox"/> Yes <input type="checkbox"/> No
	f. Has conservative treatment been attempted for at least 4 weeks/1 month?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	g. Has medication been attempted as part of conservative treatment?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	h. Has chiropractic care, physical therapy, and/or home exercise program been attempted as part of conservative treatment?	<input type="checkbox"/> Yes <input type="checkbox"/> No

<input type="checkbox"/> <b>Repeat epidural steroid injection</b> (answer a – h)	a. Is this injection being done in the same location/for the same episode of pain as the prior injection(s)?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	b. Have at least 2 weeks passed since first injection?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	c. Is the procedure planned for one caudal, one interlaminar, one bilateral transforaminal, or two unilateral transforaminal levels?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	d. Will more than the recommended amount of steroid be injected? (80mg of Triamcinolone, 80mg of methylprednisolone, 12mg of betamethasone, or 15mg of dexamethasone)	<input type="checkbox"/> Yes <input type="checkbox"/> No
	e. Does the medical record confirm at least 50% reduction in pain and improvement in function after all prior procedures for this episode of pain?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	f. Does the medical record show ongoing participation in non-operative treatment, including chiropractic/physical therapy and/or a home exercise program?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	g. Have 4 or more epidural steroid injection sessions been done in this same spine region in the past 12 months (cervical/thoracic, or lumbar)?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	h. Have 6 or more epidural steroid injection sessions for the entire spine been done in the past 6 months?	<input type="checkbox"/> Yes <input type="checkbox"/> No
<input type="checkbox"/> <b>Selective nerve root block</b> (answer a – b)	a. Is the selective nerve root block being done as a diagnostic aid to confirm which spinal level corresponds with symptoms (such as multi-level degeneration, physical exam doesn't match imaging, or confirmation of disc herniation/bulge as cause of pain)?	<input type="checkbox"/> Yes <input type="checkbox"/> No
	b. Are 2 or more levels planned for the procedure?	<input type="checkbox"/> Yes <input type="checkbox"/> No

<b>Do any of the following apply?</b> (Answer a – g)	
a. Injection with steroid planned with uncontrolled diabetes, uncontrolled hypertension, or congestive heart failure present	<input type="checkbox"/> Yes <input type="checkbox"/> No
b. Systemic or localized infection at planned injection site	<input type="checkbox"/> Yes <input type="checkbox"/> No
c. Axial neck or back pain with no or minimal radiculopathy or claudication (burning, tingling, cramping)	<input type="checkbox"/> Yes <input type="checkbox"/> No
d. Planned interlaminar ESI into insufficient epidural space (due to prior surgery, compression, or congenital condition)	<input type="checkbox"/> Yes <input type="checkbox"/> No
e. Presence of cauda equina syndrome, spinal cord compression, or spinal tumor	<input type="checkbox"/> Yes <input type="checkbox"/> No
f. Other pain management interventions planned same day (i.e. epidural steroid injection, SI joint injection, trigger point injection, etc.)	<input type="checkbox"/> Yes <input type="checkbox"/> No
g. Pain management procedures planned in multiple regions (i.e. cervical/thoracic AND lumbar or sacral)	<input type="checkbox"/> Yes <input type="checkbox"/> No
<b>Will the procedure be performed with fluoroscopic guidance?</b>	<input type="checkbox"/> Yes <input type="checkbox"/> No
<b>Is general anesthesia, conscious sedation, or monitored anesthesia care planned?</b>	<input type="checkbox"/> Yes <input type="checkbox"/> No

Include imaging reports, surgical plan and clinical documentation of all conservative therapies that have been attempted as well as the duration of each type of conservative treatment.	
<b>Form completed by:</b>	<b>Date:</b>