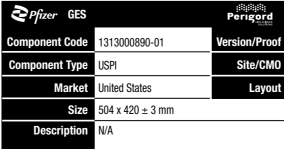
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safely and effectively. See full prescribing information for ROCURONIUM BROMIDE INJECTION.	FULL PRESCRIBING INFORMATION           1         INDICATIONS AND USAGE           Rocuronium bromide injection is indicated for inpatients and outpatients as an adjunct to general anesthesia to	2.6 Dosage in Specific Populations Pediatric Patients The recommended initial intubation dose of rocurr may be used depending on anesthetic technique a
ROCURONIUM BROMIDE injection, for intravenous use Initial U.S. Approval: 1994	facilitate both rapid sequence and routine tracheal intubation, and to provide skeletal muscle relaxation during surgery or mechanical ventilation.	For sevoflurane (induction) rocuronium bromide d
RECENT MAJOR CHANGES	2 DOSAGE AND ADMINISTRATION	to good intubating conditions within 75 seconds. Iresulted in excellent to good intubating conditions
Important Dosing and Administration Information (2.1) 07/2018 Warnings and Precautions	2.1 Important Dosing and Administration Information Rocuronium bromide injection is for intravenous use only. This drug should only be administered by	The time to maximum block for an intubating dose neonates (birth to less than 28 days). The duration
Risk of Death due to Medication Errors (5.3) 07/2018	experienced clinicians or trained individuals supervised by an experienced clinician familiar with the use, actions, characteristics, and complications of neuromuscular blocking agents. Doses of rocuronium	children (greater than 2 years up to 11 years) and lo
Rocuronium bromide injection is a nondepolarizing neuromuscular blocking agent indicated as an adjunct to general anesthesia to facilitate both rapid sequence and routine tracheal intubation, and to provide skeletal muscle relaxation during surgery or mechanical ventilation. (1)	bromide injection should be individualized and a peripheral nerve stimulator should be used to monitor drug effect, need for additional doses, adequacy of spontaneous recovery or antagonism, and to decrease the complications of overdosage if additional doses are administered.	When sevoflurane is used for induction and isoflur maintenance dosing of rocuronium bromide can b T <sub>3</sub> in all pediatric age groups. Maintenance dosing 7 to 10 mcg/kg/min, with the lowest dose requiren
To be administered only by experienced clinicians or adequately trained individuals supervised by an experienced	The dosage information which follows is derived from studies based upon units of drug per unit of body weight. It is intended to serve as an initial guide to clinicians familiar with other neuromuscular blocking agents to acquire	requirement for children (greater than 2 years up to When halothane is used for general anesthesia, pa
<ul> <li>clinician familiar with the use, actions, characteristics, and complications of neuromuscular blocking agents. (2.1)</li> <li>Individualize the dose for each patient. (2.1)</li> </ul>	experience with rocuronium bromide. In patients in whom potentiation of, or resistance to, neuromuscular block is anticipated, a dose adjustment should	administered rocuronium bromide maintenance d
<ul> <li>Peripheral nerve stimulator recommended for determination of drug response and need for additional doses, and to evaluate recovery. (2.1)</li> </ul>	be considered [see Dosage and Administration (2.6), Warnings and Precautions (5.10, 5.13), Drug Interactions (7.2, 7.3, 7.4, 7.5, 7.6, 7.8, 7.10), and Use in Specific Populations (8.6)].	clinical relaxation for 7 to 10 minutes. Alternatively of 12 mcg/kg/min upon return of T <sub>1</sub> to 10% (one tw
• Store rocuronium bromide injection with cap and ferrule intact and in a manner that minimizes the possibility	Risk of Medication Errors	neuromuscular blockade in pediatric patients. Additional information for administration to pedia
<ul> <li>of selecting the wrong product. (2.1)</li> <li><u>Tracheal intubation</u>: Recommended initial dose is 0.6 mg/kg. (2.2)</li> </ul>	Accidental administration of neuromuscular blocking agents may be fatal. Store rocuronium bromide with the cap and ferrule intact and in a manner that minimizes the possibility of selecting the wrong product [see Warnings and Precautions (5.3)].	[see Clinical Pharmacology (12.2)]. The infusion of rocuronium bromide must be indiv
<ul> <li><u>Rapid sequence intubation</u>: 0.6 to 1.2 mg/kg. (2.3)</li> <li><u>Maintenance doses</u>: Guided by response to prior dose, not administered until recovery is evident. (2.4)</li> </ul>	<b>2.2</b> Dose for Tracheal Intubation The recommended initial dose of rocuronium bromide, regardless of anesthetic technique, is 0.6 mg/kg. Neuromuscular	adjusted according to the patient's twitch response Spontaneous recovery and reversal of neuromuscu
<u>Continuous infusion</u> : Initial rate of 10 to 12 mcg/kg/min. Start only after early evidence of spontaneous recovery from an intubating dose. (2.5)	block sufficient for intubation (80% block or greater) is attained in a median (range) time of 1 (0.4 to 6) minute(s) and most patients have intubation completed within 2 minutes. Maximum blockade is achieved in most patients in less	infusion may be expected to proceed at rates com doses [see Clinical Pharmacology (12.2)].
DOSAGE FORMS AND STRENGTHS	than 3 minutes. This dose may be expected to provide 31 (15 to 85) minutes of clinical relaxation under opioid/nitrous oxide/oxygen anesthesia. Under halothane, isoflurane, and enflurane anesthesia, some extension of the period of	Rocuronium bromide is not recommended for rapi
10 mL Multiple Dose Vials containing 100 mg Rocuronium Bromide Injection (10 mg/mL) (3)CONTRAINDICATIONS	clinical relaxation should be expected [see Drug Interactions (7.3)].	Geriatric Patients Geriatric patients (65 years or older) exhibited a slig
Hypersensitivity (e.g., anaphylaxis) to rocuronium bromide or other neuromuscular blocking agents. (4)	A lower dose of rocuronium bromide (0.45 mg/kg) may be used. Neuromuscular block sufficient for intubation (80% block or greater) is attained in a median (range) time of 1.3 (0.8 to 6.2) minute(s) and most patients have intubation	62 (49 to 75), and 94 (64 to 138) minutes under opic and 1.2 mg/kg, respectively. No differences in dura
Appropriate Administration and Monitoring: Use only if facilities for intubation, mechanical ventilation, oxygen	completed within 2 minutes. Maximum blockade is achieved in most patients in less than 4 minutes. This dose may be expected to provide 22 (12 to 31) minutes of clinical relaxation under opioid/nitrous oxide/oxygen anesthesia.	rocuronium bromide were observed between thes older individuals cannot be ruled out [see Clinical P
<ul> <li>therapy, and an antagonist are immediately available. (5.1)</li> <li><u>Anaphylaxis</u>: Severe anaphylaxis has been reported. Consider cross-reactivity among neuromuscular blocking</li> </ul>	Patients receiving this low dose of 0.45 mg/kg who achieve less than 90% block (about 16% of these patients) may have a more rapid time to 25% recovery, 12 to 15 minutes.	and Precautions (5.5)]. Patients with Renal or Hepatic Impairment
agents. (5.2) <ul> <li><u>Risk of Death due to Medication Errors</u>: Accidental administration can cause death. (5.3)</li> </ul>	A large bolus dose of 0.9 or 1.2 mg/kg can be administered under opioid/nitrous oxide/oxygen anesthesia without adverse effects to the cardiovascular system [see Clinical Pharmacology (12.2)].	No differences from patients with normal hepatic a
<ul> <li><u>Need for Adequate Anesthesia</u>: Must be accompanied by adequate anesthesia or sedation. (5.4)</li> <li><u>Residual Paralysis</u>: Consider using a reversal agent in cases where residual paralysis is more likely to occur. (5.5)</li> </ul>	2.3 Rapid Sequence Intubation	0.6 mg/kg rocuronium bromide. When compared t clinical duration is similar in patients with end-stag
ADVERSE REACTIONS	In appropriately premedicated and adequately anesthetized patients, rocuronium bromide 0.6 to 1.2 mg/kg will provide excellent or good intubating conditions in most patients in less than 2 minutes [see Clinical Studies (14.1)].	1.5 times longer in patients with hepatic disease. P of effect [see Use in Specific Populations (8.6, 8.7) an
To report SUSPECTED ADVERSE REACTIONS, contact Hospira, Inc. at 1-800-441-4100 or FDA at 1-800-FDA-1088	<b>2.4 Maintenance Dosing</b> Maintenance doses of 0.1, 0.15, and 0.2 mg/kg rocuronium bromide, administered at 25% recovery of control T <sub>1</sub>	Obese Patients In obese patients, the initial dose of rocuronium br
or <u>www.fda.gov/medwatch</u> . DRUG INTERACTIONS	(defined as 3 twitches of train-of-four), provide a median (range) of 12 (2 to 31), 17 (6 to 50) and 24 (7 to 69) minutes	Weight [see Clinical Studies (14.1)]. An analysis across all US controlled clinical studies
Succinylcholine: Use before succinylcholine has not been studied. (7.11)     Nondepolarizing muscle relaxants: Interactions have been observed. (7.7)	of clinical duration under opioid/nitrous oxide/oxygen anesthesia [see Clinical Pharmacology (12.2)]. In all cases, dosing should be guided based on the clinical duration following initial dose or prior maintenance dose and not	not different between obese and non-obese patie
<ul> <li>Enhanced rocuronium bromide activity possible: Inhalation anesthetics (7.3), certain antibiotics (7.1), quinidine (7.10), magnesium (7.6), lithium (7.4), local anesthetics (7.5), procainamide. (7.8)</li> </ul>	administered until recovery of neuromuscular function is evident. A clinically insignificant cumulation of effect with repetitive maintenance dosing has been observed [see Clinical Pharmacology (12.2)].	Patients with Reduced Plasma Cholinesterase Rocuronium metabolism does not depend on plas
<u>Reduced rocuronium bromide activity possible</u> : Anticonvulsants. (7.2)	<b>2.5</b> Use by Continuous Infusion Infusion at an initial rate of 10 to 12 mcg/kg/min of rocuronium bromide should be initiated only after early evidence	patients with reduced plasma cholinesterase activity Patients with Prolonged Circulation Time
Labor and Delivery: Not recommended for rapid sequence induction in patients undergoing Cesarean section. (8.2)     Pediatric Use: Onset time and duration will vary with dose, age, and anesthetic technique. Not recommended	of spontaneous recovery from an intubating dose. Due to rapid redistribution [see Clinical Pharmacology (12.3)] and the associated rapid spontaneous recovery, initiation of the infusion after substantial return of neuromuscular function	Because higher doses of rocuronium bromide proc not be increased in these patients to reduce onset
for rapid sequence intubation in pediatric patients. (8.4)	(more than 10% of control T <sub>1</sub> ), may necessitate additional bolus doses to maintain adequate block for surgery. Upon reaching the desired level of neuromuscular block, the infusion of rocuronium bromide must be individualized	be allowed for the drug to achieve onset of effect [ Patients with Drugs or Conditions Causing Pot
See 17 for PATIENT COUNSELING INFORMATION Revised: 03/2023	for each patient. The rate of administration should be adjusted according to the patient's twitch response as monitored with the use of a peripheral nerve stimulator. In clinical trials, infusion rates have ranged from 4 to 16 mcg/kg/min.	The neuromuscular blocking action of rocuronium Potentiation is minimal when administration of the
FULL PRESCRIBING INFORMATION: CONTENTS*	Inhalation anesthetics, particularly enflurane and isoflurane, may enhance the neuromuscular blocking action of	the administration of these potent inhalation agen was 34, 38, and 42 minutes under opioid/nitrous or
1 INDICATIONS AND USAGE 2 DOSAGE AND ADMINISTRATION	nondepolarizing muscle relaxants. In the presence of steady-state concentrations of enflurane or isoflurane, it may be necessary to reduce the rate of infusion by 30% to 50%, at 45 to 60 minutes after the intubating dose.	respectively. During 1 to 2 hours of infusion, the ini 95% block was decreased by as much as 40% under
2.1 Important Dosing and Administration Information 2.2 Dose for Tracheal Intubation	Spontaneous recovery and reversal of neuromuscular blockade following discontinuation of rocuronium bromide infusion may be expected to proceed at rates comparable to that following comparable total doses administered by	2.7 Preparation for Administration of Rocurd
2.3 Rapid Sequence Intubation 2.4 Maintenance Dosing	repetitive bolus injections [see Clinical Pharmacology (12.2)]. Infusion solutions of rocuronium bromide can be prepared by mixing rocuronium bromide with an appropriate	Diluent Compatibility Rocuronium bromide is compatible in solution with
2.5 Use by Continuous Infusion 2.6 Dosage in Specific Populations	infusion solution such as 5% glucose in water or lactated Ringers [see Dosage and Administration (2.7)]. These infusion solutions should be used within 24 hours of mixing. Unused portions of infusion solutions should be discarded.	0.9% NaCl solution sterile water for 5% glucose in water lactated Ringer
2.7 Preparation for Administration of Rocuronium Bromide	Infusion rates of rocuronium bromide can be individualized for each patient using the following tables for 3 different concentrations of rocuronium bromide solution as guidelines:	5% glucose in saline Rocuronium bromide is compatible in the above so
3 DOSAGE FORMS AND STRENGTHS 4 CONTRAINDICATIONS	Table 1: Infusion Rates Using Rocuronium Bromide Injection (0.5 mg/mL)*	temperature in plastic bags, glass bottles, and plas Drug Admixture Incompatibility
5 WARNINGS AND PRECAUTIONS 5.1 Appropriate Administration and Monitoring	Patient Weight         Drug Delivery Rate (mcg/kg/min)         I           (tra)         (tra)         4         5         6         7         8         9         10         12         14         16	Rocuronium bromide is physically incompatible wi
<ul><li>5.2 Anaphylaxis</li><li>5.3 Risk of Death due to Medication Errors</li></ul>	(kg)         (lbs)         Infusion Delivery Rate (mL/hr)           10         22         4.8         6         7.2         8.4         9.6         10.8         12         14.4         16.8         19.2	amphotericin hydrocortisone sodiu amoxicillin insulin
<ul><li>5.4 Need for Adequate Anesthesia</li><li>5.5 Residual Paralysis</li></ul>	15 33 7.2 9 10.8 12.6 14.4 16.2 18 21.6 25.2 28.8	azathioprine intralipid cefazolin ketorolac
5.6 Long-Term Use in an Intensive Care Unit 5.7 Malignant Hyperthermia (MH)	20         44         9.6         12         14.4         16.8         19.2         21.6         24         28.8         33.6         38.4           25         55         12         15         18         21         24         27         30         36         42         48	cloxacillin lorazepam dexamethasone methohexital
5.8 Prolonged Circulation Time	35 77 16.8 21 25.2 29.4 33.6 37.8 42 50.4 58.8 67.2	
5.9 QT Interval Prolongation	50 110 24 30 36 42 48 54 60 72 84 96	diazepam methylprednisolone erythromycin thiopental
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> </ul>	50         110         24         30         36         42         48         54         60         72         84         96           60         132         28.8         36         43.2         50.4         57.6         64.8         72         86.4         100.8         115.2	
5.9 QT Interval Prolongation 5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block	60         132         28.8         36         43.2         50.4         57.6         64.8         72         86.4         100.8         115.2           70         154         33.6         42         50.4         58.8         67.2         75.6         84         100.8         117.6         134.4	erythromycin thiopental famotidine trimethoprim furosemide vancomycin Ilf rocuronium bromide is administered via the sam
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> </ul>	60         132         28.8         36         43.2         50.4         57.6         64.8         72         86.4         100.8         115.2           70         154         33.6         42         50.4         58.8         67.2         75.6         84         100.8         117.6         134.4           80         176         38.4         48         57.6         67.2         76.8         86.4         96         115.2         134.4         153.6           90         198         43.2         54         64.8         75.6         86.4         97.2         108         129.6         151.2         172.8	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> </ul>	60         132         28.8         36         43.2         50.4         57.6         64.8         72         86.4         100.8         115.2           70         154         33.6         42         50.4         58.8         67.2         75.6         84         100.8         117.6         134.4           80         176         38.4         48         57.6         67.2         76.8         86.4         96         115.2         134.4         153.6           90         198         43.2         54         64.8         75.6         86.4         96         115.2         134.4         153.6           100         220         48         60         72         84         96         108         120         144         168         192           *50 mg rocuronium bromide in 100 mL solution.	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed between incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> </ul>	60         132         28.8         36         43.2         50.4         57.6         64.8         72         86.4         100.8         115.2           70         154         33.6         42         50.4         58.8         67.2         75.6         84         100.8         117.6         134.4           80         176         38.4         48         57.6         67.2         76.8         86.4         96         115.2         134.4         153.6           90         198         43.2         54         64.8         75.6         86.4         96         115.2         134.4         153.6           100         220         48         60         72         84         96         108         120         144         168         192           *50 mg rocuronium bromide in 100 mL solution.           Table 2. Infusion Rates Using Rocuronium Bromide Injection (1 mg/mL)*	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> </ul>	60       132       28.8       36       43.2       50.4       57.6       64.8       72       86.4       100.8       115.2         70       154       33.6       42       50.4       58.8       67.2       75.6       84       100.8       117.6       134.4         80       176       38.4       48       57.6       67.2       76.8       86.4       96       115.2       134.4       153.6         90       198       43.2       54       64.8       75.6       86.4       97.2       108       129.6       151.2       172.8         100       220       48       60       72       84       96       108       120       144       168       192         *50 mg rocuronium bromide in 100 mL solution.         Table 2. Infusion Rates Using Rocuronium Bromide Injection (1 mg/mL)*         Patient Weight       Drug Delivery Rate (mcg/kg/min)       Image: Marcine Marci	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> <li>7.3 Inhalation Anesthetics</li> <li>7.4 Lithium Carbonate</li> </ul>	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> <li>7.3 Inhalation Anesthetics</li> <li>7.4 Lithium Carbonate</li> <li>7.5 Local Anesthetics</li> <li>7.6 Magnesium</li> </ul>	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us <b>3 DOSAGE FORMS AND STRENGTHS</b> Rocuronium bromide injection is available as:
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> <li>7.3 Inhalation Anesthetics</li> <li>7.4 Lithium Carbonate</li> <li>7.5 Local Anesthetics</li> <li>7.6 Magnesium</li> <li>7.7 Nondepolarizing Muscle Relaxants</li> <li>7.8 Procainamide</li> </ul>	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has been bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us <b>3 DOSAGE FORMS AND STRENGTHS</b>
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Table 2. Infusion Rates Using Rocuronium Bromide Injection (1 mg/mL)*           Patient Weight         Drug Delivery Rate (mcg/kg/min)         (Mg/)         (Ibs)         4         5         6         7.2         8.4         9.6           10         22         2.4         3         3.6         4.2         4.8         5.4 <t< th=""><th>erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has beer bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. 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<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Gausing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> <li>7.3 Inhalation Anesthetics</li> <li>7.4 Lithium Carbonate</li> <li>7.5 Local Anesthetics</li> <li>7.6 Magnesium</li> <li>7.7 Nordepolarizing Muscle Relaxants</li> <li>7.8 Procainamide</li> <li>7.9 Propofol</li> <li>7.10 Quinidine</li> <li>7.11 Succinylcholine</li> <li>8 USE IN SPECIFIC POPULATIONS</li> <li>8.1 Pregnancy</li> <li>8.2 Labor and Delivery</li> <li>8.4 Pediatric Use</li> <li>8.5 Geriatric Use</li> <li>8.6 Patients with Hepatic Impairment</li> <li>8.7 Patients with Renal Impairment</li> <li>8.7 Patients with Renal Impairment</li> <li>10 OVERDOSAGE</li> <li>11 DESCRIPTION</li> <li>12 CLINICAL PMRMACOLOGY</li> <li>13.1 Carcinogenesis, Intagianse, Impairment of Fertility</li> <li>14 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Pediatric Closf</li> <li>15.4 Pregramacy</li> <li>16 OVERDOSAGE</li> <li>11 DESCRIPTION</li> <li>12 CLINICAL PHARMACOLOGY</li> <li>13.1 Carcinogenesis, Impairment of Fertility</li> <li>14 Adult Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Pediatric Patients</li> </ul>	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has beer bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us <b>3 DOSAGE FORMS AND STRENGTHS</b> Rocuronium bromide injection is available as: 5 mL Multiple Dose Vials containing 50 0 10 mL Multiple Dose Vials containing 10 <b>4 CONTRAINDICATIONS</b> Rocuronium bromide is contraindicated in patients bromide or other neuromuscular blocking agents <i>f</i> <b>5 WARNINGS AND PRECAUTIONS</b> <b>5.1 Appropriate Administration and Monitor</b> Rocuronium bromide should be administered in ca experienced clinicians who are familiar with the dr drug should not be administered unless facilities for antagonist are immediately available. It is recomm agents such as rocuronium bromide employ a peri additional doses, adequacy of spontaneous recover overdosage if additional doses are administered. <b>5.2 Anaphylaxis</b> Severe anaphylactic reactions to neuromuscular bl reported. These reactions have, in some cases (ind fatal. Due to the potential severity of these reaction of appropriate emergency treatment, should be ta have had previous anaphylactic reactions to other neuromuscular blocking agents, both depolarizing <b>5.3 Risk of Death due to Medication Errors</b> Administration of rocuronium bromide results in p a progression that may be more likely to occur in
5.9       QT Interval Prolongation         5.10       Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block         5.11       Increase in Pulmonary Vascular Resistance         5.13       Use In Patients with Myasthenia         5.14       Extravasation         6       ADVERSE REACTIONS         6.1       Clinical Trials Experience         7       DRUG INTERACTIONS         7.1       Antibiotics         7.2       Anticonvulsants         7.3       Inhalation Anesthetics         7.4       Lithium Carbonate         7.5       Local Anesthetics         7.4       Lithium Carbonate         7.5       Local Anesthetics         7.6       Magnesium         7.7       Nondepolarizing Muscle Relaxants         7.8       Procainamide         7.9       Propofol         7.10       Quinidine         7.11       Succinylcholine         8       USE IN SPECIFIC POPULATIONS         8.1       Pregnancy         8.2       Labor and Delivery         8.4       Pediatric Use         8.5       Geriatric Use         8.6       Patients with Hepatic Impairment	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has beer bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us <b>3 DOSAGE FORMS AND STRENGTHS</b> Rocuronium bromide injection is available as: • 5 mL Multiple Dose Vials containing 50 • 10 mL Multiple Dose Vials containing 14 <b>4 CONTRAINDICATIONS</b> Rocuronium bromide is contraindicated in patients bromide or other neuromuscular blocking agents <i>J</i> <b>5 WARNINGS AND PRECAUTIONS</b> <b>5.1 Appropriate Administration and Monitor</b> Rocuronium bromide should be administered in ca experienced clinicians who are familiar with the dr drug should not be administered unless facilities fo antagonist are immediately available. It is recomm agents such as rocuronium bromide employ a peri additional doses, adequacy of spontaneous recove overdosage if additional doses are administered. <b>5.2 Anaphylaxis</b> Severe anaphylactic reactions to neuromuscular bl reported. These reactions have, in some cases (incl fatal. Due to the potential severity of these reaction of appropriate emergency treatment, should be ta have had previous anaphylactic reactions to other neuromuscular blocking agents, both depolarizing <b>5.3 Risk of Death due to Medication Errors</b> Administration of rocuronium bromide results in p a progression that may be more likely to occur in a of intended product and avoid confusion with othe clinical settings. If another healthcare provider is at
<ul> <li>5.9 QT Interval Prolongation</li> <li>5.10 Conditions/Drugs Gausing Potentiation of, or Resistance to, Neuromuscular Block</li> <li>5.11 Incompatibility with Alkaline Solutions</li> <li>5.12 Increase in Pulmonary Vascular Resistance</li> <li>5.13 Use In Patients with Myasthenia</li> <li>5.14 Extravasation</li> <li>6 ADVERSE REACTIONS</li> <li>6.1 Clinical Trials Experience</li> <li>6.2 Postmarketing Experience</li> <li>7 DRUG INTERACTIONS</li> <li>7.1 Antibiotics</li> <li>7.2 Anticonvulsants</li> <li>7.3 Inhalation Anesthetics</li> <li>7.4 Lithium Carbonate</li> <li>7.5 Local Anesthetics</li> <li>7.6 Magnesium</li> <li>7.7 Nondepolarizing Muscle Relaxants</li> <li>7.8 Procainamide</li> <li>7.9 Propofol</li> <li>7.10 Quinidine</li> <li>7.11 Succinylcholine</li> <li>8 USE IN SPECIFLY CPUPULATIONS</li> <li>8.1 Pregnancy</li> <li>8.2 Labor and Delivery</li> <li>8.4 Pediatric Use</li> <li>8.5 Geriatric Use</li> <li>8.6 Patients with Hepatic Impairment</li> <li>8.7 Patients with Renal Impairment of Fertility</li> <li>14 Mechanism of Action</li> <li>12.3 Pharmacodynamics</li> <li>13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility</li> <li>14 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Pediatric Datesis, Impairment of Fertility</li> <li>15 Carcinogenesis, Mutagenesis, Impairment of Fertility</li> <li>14 Adult Patients</li> <li>15 Pediatric Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Pediatric Patients</li> <li>14.3 Pediatric Patients</li> <li>15.3 Pediatric Patients</li> <li>14.3 Adult Patients</li> <li>14.3 Ad</li></ul>	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	erythromycin thiopental famotidine trimethoprim furosemide vancomycin If rocuronium bromide is administered via the sam that this infusion line is adequately flushed betwee incompatibility with rocuronium bromide has beer bromide has not been established. Infusion solutions should be used within 24 hours of discarded. Rocuronium bromide should not be mixed with all <b>Visual Inspection</b> Parenteral drug products should be inspected visu whenever solution and container permit. Do not us <b>3 DOSAGE FORMS AND STRENGTHS</b> Rocuronium bromide injection is available as: 5 mL Multiple Dose Vials containing 50 0 10 mL Multiple Dose Vials containing 10 <b>4 CONTRAINDICATIONS</b> Rocuronium bromide is contraindicated in patients bromide or other neuromuscular blocking agents <i>f</i> <b>5 WARNINGS AND PRECAUTIONS</b> <b>5.1 Appropriate Administration and Monitor</b> Rocuronium bromide should be administered in ca experienced clinicians who are familiar with the dr drug should not be administered unless facilities for antagonist are immediately available. It is recomm agents such as rocuronium bromide employ a peri additional doses, adequacy of spontaneous recover overdosage if additional doses are administered. <b>5.2 Anaphylaxis</b> Severe anaphylactic reactions to neuromuscular bl reported. These reactions have, in some cases (ind fatal. Due to the potential severity of these reaction of appropriate emergency treatment, should be ta have had previous anaphylactic reactions to other neuromuscular blocking agents, both depolarizing <b>5.3 Risk of Death due to Medication Errors</b> Administration of rocuronium bromide results in p a progression that may be more likely to occur in

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Populations	5.4 Need for Adequate Anesthesia
tubation dose of rocuronium bromide is 0.6 mg/kg; however, a lower dose of 0.45 mg/kg	Rocuronium bromide has no known effect on consciousness, pain threshold, or cerebration. Therefore, its administration must be accompanied by adequate anesthesia or sedation.
anesthetic technique and the age of the patient. rocuronium bromide doses of 0.45 mg/kg and 0.6 mg/kg in general produce excellent	5.5 Residual Paralysis In order to prevent complications resulting from residual paralysis, it is recommended to extubate only after
ons within 75 seconds. When halothane is used, a 0.6 mg/kg dose of rocuronium bromide dintubating conditions within 60 seconds.	the patient has recovered sufficiently from neuromuscular block. Geriatric patients (65 years or older) may be at
k for an intubating dose was shortest in infants (28 days up to 3 months) and longest in	increased risk for residual neuromuscular block. Other factors which could cause residual paralysis after extubation in the post-operative phase (such as drug interactions or patient condition) should also be considered. If not used
28 days). The duration of clinical relaxation following an intubating dose is shortest in rs up to 11 years) and longest in infants.	as part of standard clinical practice the use of a reversal agent should be considered, especially in those cases where residual paralysis is more likely to occur.
or induction and isoflurane/nitrous oxide for maintenance of general anesthesia, uronium bromide can be administered as bolus doses of 0.15 mg/kg at reappearance of	5.6 Long-Term Use in an Intensive Care Unit
s. Maintenance dosing can also be administered at the reappearance of $T_2$ at a rate of	Rocuronium bromide has not been studied for long-term use in the intensive care unit (ICU). As with other nondepolarizing neuromuscular blocking drugs, apparent tolerance to rocuronium bromide may develop during
he lowest dose requirement for neonates (birth to less than 28 days) and the highest dose reater than 2 years up to 11 years).	chronic administration in the ICU. While the mechanism for development of this resistance is not known, receptor up-regulation may be a contributing factor. <b>It is strongly recommended that neuromuscular transmission</b>
general anesthesia, patients ranging from 3 months old through adolescence can be romide maintenance doses of 0.075 to 0.125 mg/kg upon return of T <sub>1</sub> to 0.25% to provide	be monitored continuously during administration and recovery with the help of a nerve stimulator. Additional doses of rocuronium bromide or any other neuromuscular blocking agent should not be given
) minutes. Alternatively, a continuous infusion of rocuronium bromide initiated at a rate	until there is a definite response (one twitch of the train-of-four) to nerve stimulation. Prolonged paralysis and/or skeletal muscle weakness may be noted during initial attempts to wean from the ventilator patients who
urn of T <sub>1</sub> to 10% (one twitch present in train-of-four) may also be used to maintain pediatric patients.	have chronically received neuromuscular blocking drugs in the ICU.
administration to pediatric patients of all age groups is presented elsewhere in the label [12.2]].	Myopathy after long-term administration of other nondepolarizing neuromuscular blocking agents in the ICU alone or in combination with corticosteroid therapy has been reported. Therefore, for patients receiving both
atient's twitch response as monitored with the use of a peripheral nerve stimulator.	neuromuscular blocking agents and corticosteroids, the period of use of the neuromuscular blocking agent should be limited as much as possible and only used in the setting where in the opinion of the prescribing physician, the
reversal of neuromuscular blockade following discontinuation of rocuronium bromide	specific advantages of the drug outweigh the risk.
o proceed at rates comparable to that following similar total exposure to single bolus plogy (12.2)].	<b>5.7</b> Malignant Hyperthermia (MH) Rocuronium bromide has not been studied in MH-susceptible patients. Because rocuronium bromide is always used
recommended for rapid sequence intubation in pediatric patients.	with other agents, and the occurrence of malignant hyperthermia during anesthesia is possible even in the absence of known triggering agents, clinicians should be familiar with early signs, confirmatory diagnosis, and treatment of
or older) exhibited a slightly prolonged median (range) clinical duration of 46 (22 to 73),	malignant hyperthermia prior to the start of any anesthetic [see Adverse Reactions (6.2)].
138) minutes under opioid/nitrous oxide/oxygen anesthesia following doses of 0.6, 0.9, No differences in duration of neuromuscular blockade following maintenance doses of	In an animal study in MH-susceptible swine, the administration of rocuronium bromide injection did not appear to trigger malignant hyperthermia.
observed between these subjects and younger subjects, but greater sensitivity of some ruled out [see Clinical Pharmacology (12.2) and Clinical Studies (14.2)]. [See also Warnings	5.8 Prolonged Circulation Time Conditions associated with an increased circulatory delayed time, e.g., cardiovascular disease or advanced age, may
patic Impairment	be associated with a delay in onset time [see Dosage and Administration (2.6)].
is with normal hepatic and kidney function were observed for onset time at a dose of	<b>5.9 QT Interval Prolongation</b> The overall analysis of ECG data in pediatric patients indicates that the concomitant use of rocuronium bromide with
nide. When compared to patients with normal renal and hepatic function, the mean patients with end-stage renal disease undergoing renal transplant, and is about	general anesthetic agents can prolong the QTc interval [see Clinical Studies (14.3)].
with hepatic disease. Patients with renal failure may have a greater variation in duration Populations (8.6, 8.7) and Clinical Pharmacology (12.3)].	5.10 Conditions/Drugs Causing Potentiation of, or Resistance to, Neuromuscular Block Potentiation Nondepolarizing neuromuscular blocking agents have been found to exhibit profound neuromuscular blocking
dose of rocuronium bromide 0.6 mg/kg should be based upon the patient's actual body	effects in cachectic or debilitated patients, patients with neuromuscular diseases, and patients with carcinomatosis.
14.1)].	Certain inhalation anesthetics, particularly enflurane and isoflurane, antibiotics, magnesium salts, lithium, local anesthetics, procainamide, and quinidine have been shown to increase the duration of neuromuscular block and decrease infusion requirements of neuromuscular block and decrease infusion requirements of neuromuscular blocking agents (see Drug Interactions (7.2))
ntrolled clinical studies indicates that the pharmacodynamics of rocuronium bromide are the and non-obese patients when dosed based upon their actual body weight.	decrease infusion requirements of neuromuscular blocking agents [see Drug Interactions (7.3)]. In these or other patients in whom potentiation of neuromuscular block or difficulty with reversal may be
asma Cholinesterase Activity bes not depend on plasma cholinesterase so dosing adjustments are not needed in	anticipated, a decrease from the recommended initial dose of rocuronium bromide should be considered [see ] Dosage and Administration (2.6)].
na cholinesterase activity.	Resistance
Circulation Time uronium bromide produce a longer duration of action, the initial dosage should usually	Resistance to nondepolarizing agents, consistent with up-regulation of skeletal muscle acetylcholine receptors, is associated with burns, disuse atrophy, denervation, and direct muscle trauma. Receptor up-regulation may
atients to reduce onset time; instead, in these situations, when feasible, more time should lichieve onset of effect [see Warnings and Precautions (5.8)].	also contribute to the resistance to nondepolarizing muscle relaxants which sometimes develops in patients with cerebral palsy, patients chronically receiving anticonvulsant agents such as carbamazepine or phenytoin, or with
nditions Causing Potentiation of Neuromuscular Block g action of rocuronium bromide is potentiated by isoflurane and enflurane anesthesia.	chronic exposure to nondepolarizing agents. When rocuronium bromide is administered to these patients, shorter durations of neuromuscular block may occur, and infusion rates may be higher due to the development of resistance
en administration of the recommended dose of rocuronium bromide occurs prior to	to nondepolarizing muscle relaxants.
potent inhalation agents. The median clinical duration of a dose of 0.57 to 0.85 mg/kg under opioid/nitrous oxide/oxygen, enflurane and isoflurane maintenance anesthesia,	Severe acid-base and/or electrolyte abnormalities may potentiate or cause resistance to the neuromuscular blocking
nours of infusion, the infusion rate of rocuronium bromide required to maintain about y as much as 40% under enflurane and isoflurane anesthesia [see Drug Interactions (7.3)].	action of rocuronium bromide. No data are available in such patients and no dosing recommendations can be made. Rocuronium bromide-induced neuromuscular blockade was modified by alkalosis and acidosis in experimental
ninistration of Rocuronium Bromide	pigs. Both respiratory and metabolic acidosis prolonged the recovery time. The potency of rocuronium bromide was significantly enhanced in metabolic acidosis and alkalosis, but was reduced in respiratory alkalosis. In addition,
npatible in solution with:	experience with other drugs has suggested that acute (e.g., diarrhea) or chronic (e.g., darenocortical insufficiency) electrolyte imbalance may alter neuromuscular blockade. Since electrolyte imbalance and acid-base imbalance are
sterile water for injection lactated Ringers	usually mixed, either enhancement or inhibition may occur.
	<b>5.11</b> Incompatibility with Alkaline Solutions Rocuronium bromide, which has an acid pH, should not be mixed with alkaline solutions (e.g., barbiturate solutions)
npatible in the above solutions at concentrations up to 5 mg/mL for 24 hours at room , glass bottles, and plastic syringe pumps.	in the same syringe or administered simultaneously during intravenous infusion through the same needle.
tibility rsically incompatible when mixed with the following drugs:	<b>5.12</b> Increase in Pulmonary Vascular Resistance Rocuronium bromide may be associated with increased pulmonary vascular resistance, so caution is appropriate in
hydrocortisone sodium succinate	patients with pulmonary hypertension or valvular heart disease [see Clinical Studies (14.1)]. 5.13 Use In Patients with Myasthenia
intralipid	In patients with myasthenia gravis or myasthenic (Eaton-Lambert) syndrome, small doses of nondepolarizing
ketorolac lorazepam	neuromuscular blocking agents may have profound effects. In such patients, a peripheral nerve stimulator and use of a small test dose may be of value in monitoring the response to administration of muscle relaxants.
methohexital methylprednisolone	5.14 Extravasation
thiopental trimethoprim	If extravasation occurs, it may be associated with signs or symptoms of local irritation. The injection or infusion should be terminated immediately and restarted in another vein.
vancomycin	6 ADVERSE REACTIONS In clinical trials, the most common adverse reactions (2%) are transient hypotension and hypertension.
Iministered via the same infusion line that is also used for other drugs, it is important quately flushed between administration of rocuronium bromide and drugs for which num bromide has been demonstrated or for which compatibility with recursing the second s	The following adverse reactions are described, or described in greater detail, in other sections:
nium bromide has been demonstrated or for which compatibility with rocuronium lished.	Anaphylaxis [see Warnings and Precautions (5.2)]     Residual paralysis [see Warnings and Precautions (5.5)]
e used within 24 hours of mixing. Unused portions of infusion solutions should be	Myopathy [see Warnings and Precautions (5.6)]     Increased pulmonary vascular resistance [see Warnings and Precautions (5.12)]
d not be mixed with alkaline solutions [see Warnings and Precautions (5.11)].	6.1 Clinical Trials Experience
nould be inspected visually for particulate matter and clarity prior to administration	Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates
tainer permit. Do not use solution if particulate matter is present.	observed in practice. Clinical studies in the U.S. (n=1137) and Europe (n=1394) totaled 2531 patients. The patients exposed in the U.S.
D STRENGTHS ion is available as:	clinical studies provide the basis for calculation of adverse reaction rates. The following adverse reactions were
ose Vials containing 50 mg Rocuronium Bromide Injection (10 mg/mL) Dose Vials containing 100 mg Rocuronium Bromide Injection (10 mg/mL)	reported in patients administered rocuronium bromide (all events judged by investigators during the clinical trials to have a possible causal relationship):
IS	Adverse reactions in greater than 1% of patients: None Adverse reactions in less than 1% of patients (probably related or relationship unknown):
traindicated in patients known to have hypersensitivity (e.g., anaphylaxis) to rocuronium scular blocking agents [see Warnings and Precautions (5.2)].	Cardiovascular: arrhythmia, abnormal electrocardiogram, tachycardia Digestive: nausea, vomiting
ECAUTIONS	Respiratory: asthma (bronchospasm, wheezing, or rhonchi), hiccup
istration and Monitoring d be administered in carefully adjusted dosages by or under the supervision of	Skin and Appendages: rash, injection site edema, pruritus In the European studies, the most commonly reported reactions were transient hypotension (2%) and hypertension (2%);
are familiar with the drug's actions and the possible complications of its use. The	these are in greater frequency than the U.S. studies (0.1% and 0.1%). Changes in heart rate and blood pressure were defined differently from in the U.S. studies in which changes in cardiovascular parameters were not considered as
tered unless facilities for intubation, mechanical ventilation, oxygen therapy, and an available. It is recommended that clinicians administering neuromuscular blocking	adverse events unless judged by the investigator as unexpected, clinically significant, or thought to be histamine related. In a clinical study in patients with clinically significant cardiovascular disease undergoing coronary artery bypass
bromide employ a peripheral nerve stimulator to monitor drug effect, need for of spontaneous recovery or antagonism, and to decrease the complications of	graft, hypertension and tachycardia were reported in some patients, but these occurrences were less frequent in
ses are administered.	patients receiving beta or calcium channel-blocking drugs. In some patients, rocuronium bromide was associated with transient increases (30% or greater) in pulmonary vascular resistance. In another clinical study of patients
ns to neuromuscular blocking agents, including rocuronium bromide, have been	undergoing abdominal aortic surgery, transient increases (30% or greater) in pulmonary vascular resistance were observed in about 24% of patients receiving rocuronium bromide 0.6 or 0.9 mg/kg.
ave, in some cases (including cases with rocuronium bromide) been life threatening and everity of these reactions, the necessary precautions, such as the immediate availability	In pediatric patient studies worldwide (n=704), tachycardia occurred at an incidence of 5.3% (n=37), and it was
reatment, should be taken. Precautions should also be taken in those patients who actic reactions to other neuromuscular blocking agents, since cross-reactivity between	judged by the investigator as related in 10 cases (1.4%). 6.2 Postmarketing Experience
ents, both depolarizing and nondepolarizing, has been reported.	The following adverse reactions have been identified during post-approval use of rocuronium bromide. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably
Medication Errors Im bromide results in paralysis, which may lead to respiratory arrest and death,	estimate their frequency or establish a causal relationship to drug exposure.
nore likely to occur in a patient for whom it is not intended. Confirm proper selection oid confusion with other injectable solutions that are present in critical care and other	<i>Immune system disorders:</i> In clinical practice, there have been reports of severe allergic reactions (anaphylactic and anaphylactoid reactions and shock) with rocuronium bromide, including some that have been life-threatening and
healthcare provider is administering the product, ensure that the intended dose is clearly	fatal <i>[see Warnings and Precautions (5.2)].</i> General disorders and administration site conditions: There have been reports of malignant hyperthermia with the use
-	and the second s

of rocuronium bromide [see Warnings and Precautions (5.7)].

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## 7 DRUG INTERACTIONS

#### 7.1 Antibiotics

<sup>1</sup>Drugs which may enhance the neuromuscular blocking action of nondepolarizing agents such as rocuronium bromide include certain antibiotics (e.g., aminoglycosides; vancomycin; tetracyclines; bacitracin; polymyxins; colistin; and sodium colistimethate). If these antibiotics are used in conjunction with rocuronium bromide, prolongation of neuromuscular block may occur

## 7.2 Anticonvulsants

In 2 of 4 patients receiving chronic anticonvulsant therapy, apparent resistance to the effects of rocuronium bromide was observed in the form of diminished magnitude of neuromuscular block, or shortened clinical duration. As with other nondepolarizing neuromuscular blocking drugs, if rocuronium bromide is administered to patients chronically receiving anticonvulsant agents such as carbamazepine or phenytoin, shorter durations of neuromuscular block may loccur and infusion rates may be higher due to the development of resistance to nondepolarizing muscle relaxants. While the mechanism for development of this resistance is not known, receptor up-regulation may be a contributing factor [see Warnings and Precautions (5.10)].

7.3 Inhalation Anesthetics

Use of inhalation anesthetics has been shown to enhance the activity of other neuromuscular blocking agents

### (enflurane > isoflurane > halothane)

Isoflurane and enflurane may also prolong the duration of action of initial and maintenance doses of rocuronium bromide and decrease the average infusion requirement of rocuronium bromide by 40% compared to opioid/ nitrous oxide/oxygen anesthesia. No definite interaction between rocuronium bromide and halothane has been demonstrated. In one study, use of enflurane in 10 patients resulted in a 20% increase in mean clinical duration of the initial intubating dose, and a 37% increase in the duration of subsequent maintenance doses, when compared in

the same study to 10 patients under opioid/nitrous oxide/oxygen anesthesia. The clinical duration of initial doses of

rocuronium bromide of 0.57 to 0.85 mg/kg under enflurane or isoflurane anesthesia, as used clinically, was increased Iby 11% and 23%, respectively. The duration of maintenance doses was affected to a greater extent, increasing by 30% to 50% under either enflurane or isoflurane anesthesia. Potentiation by these agents is also observed with respect to the infusion rates of rocuronium bromide required to maintain approximately 95% neuromuscular block. Under isoflurane and enflurane anesthesia, the infusion rates are

decreased by approximately 40% compared to opioid/nitrous oxide/oxygen anesthesia. The median spontaneous recovery time (from 25% to 75% of control T<sub>1</sub>) is not affected by halothane, but is prolonged by enflurane (15% longer) and isoflurane (62% longer). Reversal-induced recovery of rocuronium bromide neuromuscular block is minimally affected by anesthetic technique [see Dosage and Administration (2.6) and Warnings and Precautions (5.10)]

#### 7.4 Lithium Carbonate

Lithium has been shown to increase the duration of neuromuscular block and decrease infusion requirements of neuromuscular blocking agents [see Warnings and Precautions (5.10)].

7.5 Local Anesthetics Local anesthetics have been shown to increase the duration of neuromuscular block and decrease infusion

requirements of neuromuscular blocking agents [ 7.6 Magnesium

#### Magnesium salts administered for the manageme [see Warninas and Precautions (5.10)].

7.7 Nondepolarizing Muscle Relaxants

There are no controlled studies documenting the use of rocuronium bromide before or after other nondepolarizing muscle relaxants. Interactions have been observed when other nondepolarizing muscle relaxants have been

## administered in succession.

7.8 Procainamide Procainamide has been shown to increase the duration of neuromuscular block and decrease infusion requirements

of neuromuscular blocking agents [see Warnings and Precautions (5.10)]. .7.9 Propofol

The use of propofol for induction and maintenance of anesthesia does not alter the clinical duration or recovery characteristics following recommended doses of rocuronium bromide.

7.10 Quinidine

Injection of quinidine during recovery from use of muscle relaxants is associated with recurrent paralysis. This possibility must also be considered for rocuronium bromide [see Warnings and Precautions (5.10)].

7.11 Succinylcholine The use of rocuronium bromide before succinylcholine, for the purpose of attenuating some of the side effects of

succinylcholine, has not been studied. If rocuronium bromide is administered following administration of succinylcholine, it should not be given until recovery from succinylcholine has been observed. The median duration of action of rocuronium bromide 0.6 mg/kg administered after a 1 mg/kg dose of succinylcholine when T<sub>1</sub> returned to 75% of control was 36 minutes (range: 14 to 57, n=12) vs. 28 minutes (range: 17 to 51, n=12) without succinylcholine.

## 8 USE IN SPECIFIC POPULATIONS

#### 8.1 Pregnancy

Developmental toxicology studies have been performed with rocuronium bromide in pregnant, conscious, nonventilated rabbits and rats. Inhibition of neuromuscular function was the endpoint for high-dose selection. The maximum tolerated dose served as the high dose and was administered intravenously 3 times a day to rats (0.3 mg/kg, 15% to 30% of human intubation dose of 0.6 to 1.2 mg/kg based on the body surface unit of mg/m<sup>2</sup>) from Day 6 to 17 and to rabbits (0.02 mg/kg, 25% human dose) from Day 6 to 18 of pregnancy. High-dose treatment caused acute symptoms of respiratory dysfunction due to the pharmacological activity of the drug. Teratogenicity was not observed in these animal species. The incidence of late embryonic death was increased at the high dose in rats, most likely due to oxygen deficiency. Therefore, this finding probably has no relevance for humans because immediate mechanical ventilation of the intubated patient will reffectively prevent embryo-fetal hypoxia. However, there are no adequate and well-controlled studies in pregnant women.

#### Rocuronium bromide should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. .8.2 Labor and Delivery

The use of rocuronium bromide in Cesarean section has been studied in a limited number of patients [see Clinical Studies (14.1)]. Rocuronium bromide is not recommended for rapid sequence induction in Cesarean section patients.

8.4 Pediatric Use ide has been studied in pediatric patients 3 months to 14 years of age under halothane anesthesia. Of the pediatric patients anesthetized with halothane who did not receive atropine for induction, about 80% experienced a transient increase (30% or greater) in heart rate after intubation. One of the 19 infants anesthetized with halothane and fentanyl who received atropine for induction experienced this magnitude of change [see Dosage and Administration (2.6) and Clinical Studies (14.3)].

Rocuronium bromide was also studied in pediatric patients up to 17 years of age, including neonates, under sevoflurane (induction) and isoflurane/nitrous oxide (maintenance) anesthesia. Onset time and clinical duration varied with dose, the age of the patient, and anesthetic technique.

The overall analysis of ECG data in pediatric patients indicates that the concomitant use of rocuronium bromide with general anesthetic agents can prolong the QTc interval. The data also suggest that rocuronium bromide may increase heart rate. However, it was not possible to conclusively identify an effect of rocuronium bromide independent of that of anesthesia and other factors. Additionally, when examining plasma levels of rocuronium bromide in correlation to QTc interval prolongation, no relationship was observed [see Dosage and Administration] (2.6), Warnings and Precautions (5.9), and Clinical Studies (14.3)].

Rocuronium bromide is not recommended for rapid sequence intubation in pediatric patients. Recommendations for use in pediatric patients are discussed in other sections [see Dosage and Administration (2.6) and Clinical Pharmacology (12.2)]. 8.5 Geriatric Use

Rocuronium bromide was administered to 140 geriatric patients (65 years or greater) in U.S. clinical trials and 128 geriatric patients in European clinical trials. The observed pharmacokinetic profile for geriatric patients (n=20) was similar to that for other adult surgical patients [see Clinical Pharmacology (12.3)]. Onset time and duration of action were slightly longer for geriatric patients (n=43) in clinical trials. Clinical experiences and recommendations for use in geriatric patients are discussed in other sections [see Dosage and Administration (2.6), Warnings and Precautions (5.5), Clinical Pharmacology (12.2), and Clinical Studies (14.2)].

8.6 Patients with Hepatic Impairment Since rocuronium bromide is primarily excreted by the liver, it should be used with caution in patients with clinically significant hepatic impairment. Rocuronium bromide 0.6 mg/kg has been studied in a limited number of patients (n=9) with clinically significant hepatic impairment under steady-state isoflurane anesthesia. After rocuronium bromide 10.6 mg/kg, the median (range) clinical duration of 60 (35 to 166) minutes was moderately prolonged compared to 42 minutes in patients with normal hepatic function. The median recovery time of 53 minutes was also prolonged in patients with cirrhosis compared to 20 minutes in patients with normal hepatic function. Four of 8 patients with cirrhosis, who received rocuronium bromide 0.6 mg/kg under opioid/nitrous oxide/oxygen anesthesia, did not achieve

complete block. These findings are consistent with the increase in volume of distribution at steady state observed in patients with significant hepatic impairment [see Clinical Pharmacology (12.3)]. If used for rapid sequence induction in patients with ascites, an increased initial dosage may be necessary to assure complete block. Duration will be prolonged in these cases. The use of doses higher than 0.6 mg/kg has not been studied [see Dosage and Administration (2.6)]. 8.7 Patients with Renal Impairment

Due to the limited role of the kidney in the excretion of rocuronium bromide, usual dosing guidelines should be followed. In patients with renal dysfunction, the duration of neuromuscular blockade was not prolonged: however, there was substantial individual variability (range: 22 to 90 minutes) [see Clinical Pharmacology (12.3)]. 10 OVERDOSAGE

Overdosage with neuromuscular blocking agents may result in neuromuscular block beyond the time needed for surgery and anesthesia. The primary treatment is maintenance of a patent airway, controlled ventilation, and adequate sedation until recovery of normal neuromuscular function is assured. Once evidence of recovery from

neuromuscular block is observed, further recovery may be facilitated by administration of an anticholinesterase agent in conjunction with an appropriate anticholinergic agent. Reversal of Neuromuscular Blockade: Anticholinesterase agents should not be administered prior to the demonstration of some spontaneous recovery from neuromuscular blockade. The use of a nerve stimulator to document recovery is recommended. Patients should be evaluated for adequate clinical evidence of neuromuscular recovery, e.g., 5-second head lift, adequate phonation, ventilation, and upper airway patency. Ventilation must be supported while patients exhibit any signs of muscle weakness. Recovery may be delayed in the presence of debilitation, carcinomatosis, and concomitant use of certain drugs

which enhance neuromuscular blockade or separately cause respiratory depression. Under such circumstances the

## management is the same as that of prolonged neuromuscular blockade.

The structural formula is:

11 DESCRIPTION Rocuronium bromide injection is a nondepolarizing neuromuscular blocking agent with a rapid to intermediate onset depending on dose and intermediate duration. Rocuronium bromide is chemically designated as 1-[17β-(acetyloxy)-3α-hydroxy-2β-(4-morpholinyl)-5α-androstan-16β-yl]-1-(2-propenyl)pyrrolidinium bromide.

Br<sup>\*</sup>

The chemical formula is C<sub>32</sub>H<sub>53</sub>BrN<sub>2</sub>O<sub>4</sub> with a molecular weight of 609.70. The partition coefficient of rocuronium bromide in n-octanol/water is 0.5 at 20°C

Rocuronium bromide is supplied as a sterile, nonpyrogenic, isotonic solution that is clear, colorless to yellow/orange, for intravenous injection only. Each mL contains 10 mg rocuronium bromide and 2 mg sodium acetate. The aqueous solution is adjusted to isotonicity with sodium chloride and to a pH of 4 with acetic acid and/or sodium hydroxide. 12 CLINICAL PHARMACOLOGY

#### 12.1 Mechanism of Action

Rocuronium bromide is a nondepolarizing neuromuscular blocking agent with a rapid to intermediate onset depending on dose and intermediate duration. It acts by competing for cholinergic receptors at the motor end-plate. This action is antagonized by acetylcholinesterase inhibitors, such as neostigmine and edrophonium.

12.2 Pharmacodynamics The ED<sub>95</sub> (dose required to produce 95% suppression of the first [T<sub>1</sub>] mechanomyographic [MMG] response of the adductor pollicis muscle [thumb] to indirect supramaximal train-of-four stimulation of the ulnar nerve) during opioid/nitrous oxide/oxygen anesthesia is approximately 0.3 mg/kg. Patient variability around the ED<sub>95</sub> dose sts that 50% of patients will exhibit  $T_1$  depression of 91% to 97%.

Table 4 presents intubating condition	ns in patients with intubation initiated at 6	0 to 70 seconds.
	lent or Good Intubating Conditions an ation in Patients with Intubation Initiat	
Rocuronium Bromide Dose (mg/kg) Administered over 5 sec	Percent of Patients with Excellent or Good Intubating Conditions	Time to Completion of Intubation (min)
Adult*18 to 64 yrs 0.45 (n=43) 0.6 (n=51)	86% 96%	1.6 (1.0 to 7.0) 1.6 (1.0 to 3.2)
Infants <sup>+</sup> 3 mo to 1 yr 0.6 (n=18) Pediatric <sup>+</sup> 1 to 12 yrs	100%	1.0 (1.0 to 1.5)
0.6 (n=12)	100%	1.0 (0.5 to 2.3)

\* Excludes patients undergoing Cesarean section. Pediatric patients were under halothane anesthesia.

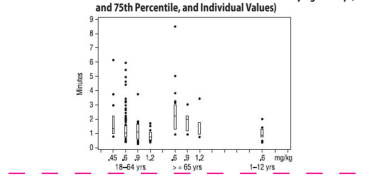
Excellent intubating conditions=jaw relaxed, vocal cords apart and immobile, no diaphragmatic movement. Good intubating conditions=same as excellent but with some diaphragmatic movement. Table 5 presents the time to onset and clinical duration for the initial dose of rocuronium bromide injection under opioid/

Table 5: Median (Range) Time to Onse Opioid/Nitrous Oxide/Oxygen Ane		alothane Ánesthesia (F	Pediatric Patients)
Rocuronium Bromide Dose (mg/kg) Administered over 5 sec	Time to ≥80% Block (min)	Time to Maximum Block (min)	Clinical Duration (min)
Adult 18 to 64 yrs 0.45 (n=50) 0.6 (n=142) 0.9 (n=20) 1.2 (n=18)	1.3 (0.8 to 6.2) 1.0 (0.4 to 6.0) 1.1 (0.3 to 3.8) 0.7 (0.4 to 1.7)	3.0 (1.3 to 8.2) 1.8 (0.6 to 13.0) 1.4 (0.8 to 6.2) 1.0 (0.6 to 4.7)	22 (12 to 31) 31 (15 to 85) 58 (27 to 111) 67 (38 to 160)
Geriatric ≥65 yrs 0.6 (n=31) 0.9 (n=5) 1.2 (n=7)	2.3 (1.0 to 8.3) 2.0 (1.0 to 3.0) 1.0 (0.8 to 3.5)	3.7 (1.3 to 11.3) 2.5 (1.2 to 5.0) 1.3 (1.2 to 4.7)	46 (22 to 73) 62 (49 to 75) 94 (64 to 138)
Infants 3 mo to 1 yr 0.6 (n=17) 0.8 (n=9)	-	0.8 (0.3 to 3.0) 0.7 (0.5 to 0.8)	41 (24 to 68) 40 (27 to 70)
Pediatric 1 to 12 yrs 0.6 (n=27) 0.8 (n=18)	0.8 (0.4 to 2.0)	1.0 (0.5 to 3.3) 0.5 (0.3 to 1.0)	26 (17 to 39) 30 (17 to 56)

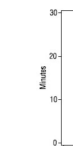
Clinical duration=time until return to 25% of control T1. Patients receiving doses of 0.45 mg/kg who achieved less than 90% block (16% of these patients) had about 12 to 15 minutes to 25% recovery. Table 6 presents the time to onset and clinical duration for the initial dose of rocuronium bromide injection under Isevoflurane (induction) and isoflurane/nitrous oxide (maintenance) anesthesia in pediatric patients.

Rocuronium Bromide Dose (mg/kg) Administered over 5 sec	Time to Maximum Block (min)	Time to Reappearance T (min)
Neonates birth to <28 days		
0.45 (n=5)	1.1 (0.6 to 2.2)	40.3 (32.5 to 62.6)
0.6 (n=10)	1.0 (0.2 to 2.1)	49.7 (16.6 to 119.0)
1 (n=6)	0.6 (0.3 to 1.8)	114.4 (92.6 to 136.3)
Infants 28 days to ≤3 mo		
0.45 (n=9)	0.5 (0.4 to 1.3)	49.1 (13.5 to 79.9)
0.6 (n=11)	0.4 (0.2 to 0.8)	59.8 (32.3 to 87.8)
1 (n=5)	0.3 (0.2 to 0.7)	103.3 (90.8 to 155.4)
Toddlers >3 mo to ≤2 yrs		
0.45 (n=17)	0.8 (0.3 to 1.9)	39.2 (16.9 to 59.4)
0.6 (n=29)	0.6 (0.2 to 1.6)	44.2 (18.9 to 68.8)
1 (n=15)	0.5 (0.2 to 1.5)	72.0 (36.2 to 128.2)
Children >2 yrs to ≤11 yrs		
0.45 (n=14)	0.9 (0.4 to 1.9)	21.5 (17.5 to 38.0)
0.6 (n=37)	0.8 (0.3 to 1.7)	36.7 (20.1 to 65.9)
1 (n=16)	0.7 (0.4 to 1.2)	53.1 (31.2 to 89.9)
Adolescents >11 to ≤17 yrs		
0.45 (n=18)	1.0 (0.5 to 1.7)	37.5 (18.3 to 65.7)
0.6 (n=31)	0.9 (0.2 to 2.1)	41.4 (16.3 to 91.2)
1 (n=14)	0.7 (0.5 to 1.2)	67.1 (25.6 to 93.8)

The time to 80% or greater block and clinical duration as a function of dose are presented in Figures 1 and 2. Figure 1: Time to 80% or Greater Block vs. Initial Dose of Rocuronium Bromide by Age Group (Median, 25th



represented in Figure 3 [see Dosage and Administration (2.4)].



Once spontaneous recovery has reached 25% of control T<sub>1</sub>, the neuromuscular block produced by rocuronium bromide is readily reversed with anticholinesterase agents, e.g., edrophonium or neostigmine The median spontaneous recovery from 25% to 75% T<sub>1</sub> was 13 minutes in adult patients. When neuromuscular block was reversed in 36 adults at a  $T_1$  of 22% to 27%, recovery to a  $T_1$  of 89 (50 to 132)% and  $T_4/T_1$  of 69 (38 to 92)% was achieved within 5 minutes. Only 5 of 320 adults reversed received an additional dose of reversal agent. The median (range) dose of neostigmine was 0.04 (0.01 to 0.09) mg/kg and the median (range) dose of edrophonium was

0.5 (0.3 to 1.0) ma/ka In geriatric patients (n=51) reversed with neostigmine, the median  $T_4/T_1$  increased from 40% to 88% in 5 minutes. In clinical trials with halothane, pediatric patients (n=27) who received 0.5 mg/kg edrophonium had increases in the median T<sub>4</sub>/T<sub>1</sub> from 37% at reversal to 93% after 2 minutes. Pediatric patients (n=58) who received 1 mg/kg edrophonium had increases in the median  $T_4/T_1$  from 72% at reversal to 100% after 2 minutes. Infants (n=10) who were reversed with 0.03 mg/kg neostigmine recovered from 25% to 75% T<sub>1</sub> within 4 minutes. There were no reports of less than satisfactory clinical recovery of neuromuscular function.

anesthetics [see Drug Interactions (7.3)]. Hemodynamics

There were no dose-related effects on the incidence of changes from baseline (30% or greater) in mean arterial blood pressure (MAP) or heart rate associated with rocuronium bromide administration over the dose range of 0.12 to 1.2 mg/kg (4 × ED<sub>95</sub>) within 5 minutes after rocuronium bromide administration and prior to intubation. Increases or decreases in MAP were observed in 2% to 5% of geriatric and other adult patients, and in about 1% of pediatric patients. Heart rate changes (30% or greater) occurred in 0% to 2% of geriatric and other adult patients. Tachycardia (30% or greater) occurred in 12 of 127 pediatric patients. Most of the pediatric patients developing tachycardia were from a single study where the patients were anesthetized with halothane and who did not receive atropine for induction [see Clinical Studies (14.3)]. In U.S. studies, laryngoscopy and tracheal intubation following rocuronium bromide administration were accompanied by transient tachycardia (30% or greater increases) in about one-third of adult patients under opioid/nitrous oxide/oxygen anesthesia. Animal studies have indicated that the ratio of vagal:neuromuscular block following rocuronium bromide administration is less than vecuronium but Igreater than pancuronium. The tachycardia observed in some patients may result from this vagal blocking activity.

Histamine Release

12.3 Pharmacokinetics Adult and Geriatric Patients

variability are contained in the following section.

## PK Parameters

Clearance (L/kg/hr) Volume of Distribution at Steady Stat  $t_{1/2}\beta$  Elimination (hr)

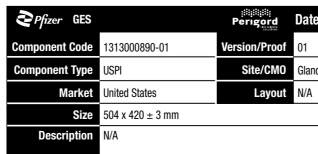
In general, studies with normal adult subjects did not reveal any differences in the pha due to gender. pharmacodynamics of rocuronium in humans are consistent with these findings.

#### **PK Parameters** Clearance (L/kg/hr) Volume of Distribution at Steady Stat

 $_{1/2} \beta$  Elimination (hr) \*Differences in the calculated  $t_{1/2}\beta$  and Cl between this study and the study in young adults vs. geriatrics ( $\geq$ 65 years) is related to the different sample populations and anesthetic techniques. The net result of these findings is that subjects with renal failure have clinical durations that are similar to but somewhat more variable than the duration that one would expect in subjects with normal renal function. Hepatically impaired patients, due to the large increase in volume, may demonstrate clinical durations approaching 1.5 times that of subjects with normal hepatic function. In both populations the clinician should individualize the dose to the needs of the patient [see Dosage and Administration (2.6)]. Tissue redistribution accounts for most (about 80%) of the initial amount of rocuronium administered. As tissue compartments fill with continued dosing (4 to 8 hours), less drug is redistributed away from the site of action and, for an infusion-only dose, the rate to maintain neuromuscular blockade falls to about 20% of the initial infusion rate. The use of a loading dose and a smaller infusion rate reduces the need for adjustment of dose. Pediatric Patients

Under halothane anesthesia, the clinical duration of effects of rocuronium bromide did not vary with age in patients 4 months to 8 years of age. The terminal half-life and other pharmacokinetic parameters of rocuronium in these pediatric patients are presented in Table 9.

## 504 mm



[see Warnings and Precautions (5.10)].	suggest Table 4
ent of toxemia of pregnancy may enhance neuromuscular blockade	

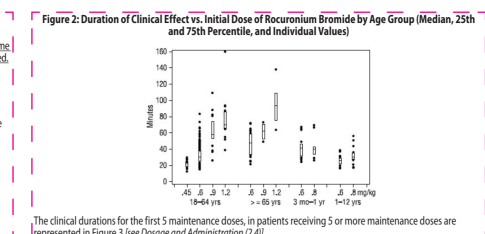
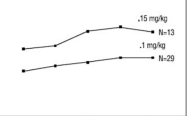


Figure 3: Duration of Clinical Effect vs. Number of Rocuronium Bromide Maintenance Doses, by Dose



The neuromuscular blocking action of rocuronium bromide may be enhanced in the presence of potent inhalation

In studies of histamine release, clinically significant concentrations of plasma histamine occurred in 1 of 88 patients. Clinical signs of histamine release (flushing, rash, or bronchospasm) associated with the administration of rocuronium bromide were assessed in clinical trials and reported in 9 of 1137 (0.8%) patients.

In an effort to maximize the information gathered in the *in vivo* pharmacokinetic studies, the data from the studies was used to develop population estimates of the parameters for the subpopulations represented (e.g., geriatric, pediatric, renal, and hepatic impairment). These population-based estimates and a measure of the estimate

Following intravenous administration of rocuronium bromide, plasma levels of rocuronium follow a threecompartment open model. The rapid distribution half-life is 1 to 2 minutes and the slower distribution half-life is 14 to 18 minutes. Rocuronium is approximately 30% bound to human plasma proteins. In geriatric and other adult surgical patients undergoing either opioid/nitrous oxide/oxygen or inhalational anesthesia, the observed pharmacokinetic profile was essentially unchanged [see Dosage and Administration (2.6)].

Table 7: Mean (SD) Pharmacokinetic Parameters in Adults (n=22; ages 27 to 58 yrs) and Geriatric (n=20;

65 yrs or greater) During Opioid/Nitrous Oxide/Oxygen Anesthesia		
	Adults (Ages 27 to 58 yrs)	Geriatrics (≥65 yrs)
	0.25 (0.08)	0.21 (0.06)
tion at Steady State (L/kg)	0.25 (0.04)	0.22 (0.03)
()	1.4 (0.4)	1.5 (0.4)
vith normal adult subjects did no	t reveal any differences in the pl	parmacokinetics of rocuronium

Studies of distribution, metabolism, and excretion in cats and dogs indicate that rocuronium is eliminated primarily by the liver. The rocuronium analog 17-desacetyl-rocuronium, a metabolite, has been rarely observed in the plasma prurine of humans administered single doses of 0.5 to 1 mg/kg with or without a subsequent infusion (for up to 12 hr) of rocuronium. In the cat, 17-desacetyl-rocuronium has approximately one-twentieth the neuromuscular blocking potency of rocuronium. The effects of renal failure and hepatic disease on the pharmacokinetics and

In general, patients undergoing cadaver kidney transplant have a small reduction in clearance which is offset pharmacokinetically by a corresponding increase in volume, such that the net effect is an unchanged plasma half-life. Patients with demonstrated liver cirrhosis have a marked increase in their volume of distribution resulting

in a plasma half-life approximately twice that of patients with normal hepatic function. Table 8 shows the pharmacokinetic parameters in subjects with either impaired renal or hepatic function. Table 8: Mean (SD) Pharmacokinetic Parameters in Adults with Normal Renal and Hepatic Function (n=10,

ages 23 to 65), Renal Transplant Patients (n=10, ages 21 to 45), and Hepatic Dysfunction Patients (n=9, ages 31 to 67) During Isoflurane Anesthesia Normal Renal and Renal Transplant Hepatic Dysfunction

	Hepatic Function	Patients	Patients
	0.16 (0.05)*	0.13 (0.04)	0.13 (0.06)
te (L/kg)	0.26 (0.03)	0.34 (0.11)	0.53 (0.14)
	2.4 (0.8)*	2.4 (1.1)	4.3 (2.6)

PK Parameters Patient Age Range					
Clearance (L/kg/hr)			1 to <3 yrs		
Volume of Distribution at State (L/kg)	Steady	0.30 (0.04)	0.26 (0.06)		0.21 (0.03)
$t_{1/2}\beta$ Elimination (hr)		1.3 (0.5) 1.1 (0.7) 0.8 (0.3)		0.8 (0.3)	
Pharmacokinetics of rocur datasets from 2 trials under oharmacokinetic paramet 18 years clearance (CL) and the terminal half-life of ro- 10 presents the pharmaco and isoflurane/nitrous oxi <b>Table 10: Mean (SD) Ph</b> (ind	er sevoflurane (in ters were found to d volume of distri curonium bromic okinetic paramete de (maintenance <b>armacokinetic F</b>	Iduction) and isoflurar o be linearly proportic ibution (Vss) increase de decreases with incr ers in the different age ) anesthesia. Parameters of Rocur flurane/Nitrous Oxic	ne/nitrous oxide (n mal to body weigh with bodyweight easing age from 1. groups in the stur onium in Pediatr le (maintenance)	naintenance) an ht. In patients u (kg) and age (ye hour to 0.7 to dies with sevof ic Patients Du	nesthesia. All nder the age of ears). As a result 0.8 hour. Table lurane (induction)
PK Parameters	Birth to <28 day	Pat s 28 days to ≤3 mos	ient Age Range 3 mos to ≤2 yrs	2 to ≤11 yrs	11 to ≤17 yrs
CL (L/kg/hr) Volume of Distribution	0.31 (0.07)	0.30 (0.08)	0.33 (0.10)	0.35 (0.09)	0.29 (0.14)
(L/kg)	0.42 (0.06)	0.31 (0.03)	0.23 (0.03)	0.18 (0.02)	0.18 (0.01)
t <sub>1/2</sub> β (hr)	1.1 (0.2)	0.9 (0.3)	0.8 (0.2)	0.7 (0.2)	0.8 (0.3)
In U.S. clinical studies, a too 140 geriatric, 55 obstetric, ASA III, and 10 patients (ur clinical studies, a total of 1 (65 years or greater), and 1 <b>14.1 Adult Patients</b> Intubation using doses of Excellent to good intubatii within 3 minutes in most p 33 (14 to 85) minutes unded in 2 studies with 19 and 16 67 (38 to 160) minutes of of <b>Cardiovascular Disease</b> In 1 clinical study, 10 patie graft received an initial do surgery with bolus mainte rocuronium bromide proc surgical intensive care uni <b>Rapid Sequence Intubati</b> 6 mg/kg) or propofol (1.5 m Most of the patients also r attempted within 60 to 90 (1.5 mg/kg. Excellent or go 95% to 99.9%]) patients succinylcholine. The durat dose is approximately equ <b>Obese Patients</b> Rocuronium bromide in weight (IBW) was not asso for cruonium bromide in weight (IBW) was not asso for rapid sequence induct 1.1. Obese patients dos clinical duration of 25 (14 i based on ABW. These resu weight [ <i>see Dosage and Ac</i> <b>Obstetric Patients</b> Rocuronium bromide 0.6 of for rapid sequence induct 5 minutes. The umbilical v conditions were poor or ir attempted 60 seconds aft induction in Cesarean secci <b>14.3 Pediatric Patients</b> Rocuronium bromide vase 0.6 mg/kg vas shortest in dose in approving provided excell Recovery times from 25% patients [ <i>see Dosage and Ac</i> <b>14.3 Pediatric Patients</b> Rocuronium bromide vase and infusion requirement: maximum block in about 0.6 mg/kg was shortest in	and 766 other ac ndergoing corona 1394 patients rece 1214 other adults rocuronium brom ing conditions we patients. Dosew of opioid/nitrous 5 patients under of clinical relaxation ents with clinically ose of 0.6 mg/kg i enance doses of 0 duced relaxation it (SICU) while the <b>tion</b> n patients in 6 clii to 2.5 mg/kg) in of received a preme 0 seconds of adm ood intubating co ceiving rocuroni tion of action of ru- uivalent to the du s dosed according the 47 of 330 (14 octated with clinical uced neuromuscu e patients, rocuro sed according to to 2.9) minutes, an ults support the re dministration (2.6) mg/kg was admini ion of an esthesia renous plasma co hadequate in 5 of er drug injection. tion patients. s evaluated in 55 s lent 75% after thes ddministration (2.4) f, 0.6, or 1 mg/kg thesia for intubat to xere evaluated 1 minute. Across the children [36.	dults. Most patients (9 ary artery bypass grafi eived rocuronium brom- mide 0.6 to 0.85 mg/kg ere generally achieved vithin this range provice oxide/oxygen anesthe oppioid/nitrous oxide/oc, respectively. y significant cardiovas rocuronium bromide. .3 mg/kg. Following in sufficient to support ne e patients were recove nical studies where an combination with eith dication such as mida- inistration of rocuroni nditions were achieve um bromide and in 10 ocuronium bromide 0 ration of other interm g to actual body weigh %) patients who were cally significant differe ular block. nium bromide 0.6 mg IBW had a longer time nd did not achieve intu ecommendation that of <i>J</i> . mistered with thiopent for Cesarean section. incentrations were 18 <sup>6</sup> 13 women receiving 3 . Therefore, rocuronium geriatric patients (age bating conditions in a se doses were not prol <i>6) and Use in Specific Pi</i> was evaluated under in 137 patients. In all all age groups, media 7 (20.1 to 65.9) minute	0%) were ASÁ phy ing or valvular sum mide injection, inc g was evaluated in within 2 minutes de clinical relaxatic sia. Larger doses ( xygen anesthesia cular disease unde Neuromuscular blin nduction, continuu nechanical ventilar ring from surgery. esthesia was indu er fentanyl (2 to 5 zolam or temazep um bromide 0.6 m d in 119/120 (99% 8/110 (98% [94% t .6 mg/kg is longer ediate-acting neu ti (ABW) in most cl at least 30% or m nnces in the onset, /kg was dosed acc to maximum blog base patients be stal, 3 to 4 mg/kg (r No neonate had A % of maternal com 3 to 4 mg/kg thiop m bromide is not r s 65 to 80 years) in median (range) tim onged in geriatric opulations (8.5)].	sical status for gery) were AS/ luding 52 pedia 203 adults in 1 and maximum on for a mediam 0.9 and 1.2 mg and provided 5 ergoing corona ock was mainta ous 8 mcg/kg/r tion for 6 to 12 ced with either mcg/kg) or alfe am. Most patie tg/kg or succiny [95% confiden o 99.8%]) patie than succinylc romuscular blo inical studies. 1 ore above their duration, recov ording to ABW ck, a shorter me s comparable to dosed based o a=13) or 4 to 6 r PGAR scores gi centrations at c ental when int ecommended 6 clinical studie for a studies fun associations at c ental when int ecommended 6 clinical studies fun associations at c ental studies f	II, about 9% were AIV. In European atric, 128 geriatric 1 clinical studies. block occurred (range) time of /kg) were evaluated is (27 to 111) and ry artery bypass ined during nin infusion of hours in the thiopental (3 to intanil (1 mg). nts had intubation ylcholine 1 to ce interval: ints receiving holine and at this cking drugs. The administration i deal body very, or reversal of (n=12) or IBW edian (range) o those dosed n actual body mg/kg (n=42), reater than 7 at lelivery. Intubating ubation was for rapid sequence es. Doses of 3) minutes. ared to other adult urane/nitrous aintenance bolus rovided time to fT <sub>3</sub> for doses of .3 to 87.8) minutes]
For pediatric patients olde when compared with bolu Rocuronium bromide 0.6 of 12 months, n=47; age 1 to oxygen. Doses of 0.6 mg/l provided a median (range 26 (17 to 39) minutes in 1- Populations (8.4)]. <b>16 HOW SUPPLIED/ST</b>	er than 3 months, us maintenance [: or 0.8 mg/kg was o 12 years) in 3 stu kg provided a me o) time of clinical of to 12-year-old pe <b>TORAGE AND HA</b>	, the time to recovery see Dosage and Admin sevaluated for intubat idies using halothane idian (range) time to n relaxation of 41 (24 to ediatric patients [see D INDLING	was shorter after s istration (2.6) and l ion in 75 pediatric (1% to 5%) and nit naximum block of 68) minutes in 3-n	topping infusio <i>Jse in Specific P</i> patients (n=28 rous oxide (609 1 (0.5 to 3.3) m nonth to 1-year	on maintenance opulations (8.4)]. b; age 3 to % to 70%) in inute(s). This dose -old infants and
0409-3189-10 50 mg/5 0409-7037-10 100 mg/ <b>Storage Conditions</b>	<b>nium Bromide I</b> 5 mL Multiple Dos /10 mL Multiple D	<b>njection (10 mg/mL)</b> se Vial Dose Vial	10 vials per 10 vials per	carton carton	
J	ction should be s n to room tempe of rocuronium b	tored in a refrigerator rature storage conditi romide within 30 days	, 2° to 8°C (36° to 4 ons (25°C/77°F), u:	6°F). <b>Do not fr</b> se rocuronium	bromide within
17 PATIENT COUNSEL Obtain information about rocuronium bromide or of medical conditions and m In addition, inform your par rocuronium bromide, have information from your par	ING INFORMATI your patient's m ther neuromuscu redications might atient that severe e been reported.	edical history, current lar blocking agents. If influence how rocurc anaphylactic reaction Since allergic cross-re	applicable, inform nium bromide wo ns to neuromuscul activity has been r	n your patients orks. ar blocking age reported in this	that certain ents, including class, request

Non-Print Colors Perigord Date: 20 Jun 2023 Print Colors Time: 17:36 Black Profile PAR Number PAR-2023-0004392 Component Code OLD Gland Pharma 1313000890-00 Site Technical Code N/A Smallest BODY TEXT size

# 420 mm