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14. TRANSPORT INFORMATION		
UN Number:	Not available	
UN Proper	Not regulated for transport of dangerous goods	
Shipping Name:		
Transport Hazard	Non-dangerous goods	
Class(es):		
Packing Group:	None	
Environmental	Non-hazardous	
Hazards:		
Special	None	
Requirements:		

15. REGULATORY INFORMATION	
Other Regulatory	EC Classification: Exempt.
Information:	
	TSCA (Toxic Substances Control Act) Regulations, 40CFR 710: This product is a drug and is exempt from TSCA regulation.
	OSHA Status: Hazardous as defined by OSHA 29 CFR 1910. 1200 (c)
	CERCLA and SARA Regulations (40CFR 355,370 and 372): This product does not contain any chemical subject to the reporting requirements of SARA Section 313.
	Federal Regulatory Information: Regulated under OSHA and FDA
	State Regulatory Information: Consult with state environmental and/or public health agencies.
	Health Canada: Class D2 (Materials causing other toxic effects)

16. OTHER INFORMATION	
Date of Preparation	10 July 2013
of this MSDS:	
References:	Toxicological information obtained from RTECS, Toxline and publicly available sources

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12. ECOLOGIC	CAL INFORMATION	
No information on this formulation. The product is soluble in water. The following		
information refers to active ingredient prilocaine:		
Toxicity:	Harmful to aquatic organisms. LC50 (zebra fish) (96 hour)	
	188 mg/L EC50 (Daphnia magna) (48 hour) 61 mg/L. EC50	
	(green algae) (72 hour) 154 mg/L.	
Persistence and	May cause long-term adverse effects in the aquatic	
Degradability:	environment. Not readily biodegradable. (ISO7827-1984(E))	
Bioaccumulation	Log Kow:	
Potential:	Lidocaine base: 2.44 (experimental)	
	Prilocaine base: 2.11	
	Log Koc:	
	Lidocaine base: 2.623 (MCI method)	
	Prilocaine base: 2.611 (MCI method)	
	Log BCF:	
	Lidocaine base: 1.277	
	Prilocaine base: 1.059	
	Atmospheric Half-Life:	
	Lidocaine base: 2.35 h	
	Prilocaine base: 2.46 h	
	as calculated via applicable algorithms encoded in EpiSuite 4.0	
	(USEPA 2010)	
Mobility in Soil:	No information available	

13. DISPOSAL CONSIDERATIONS		
Disposal Methods:	Dissolve or mix the material with a combustible solvent and burn in a chemical incinerator equipped with an afterburner and scrubber. Observe all federal, state and local environmental regulations.	

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	caution exercised when administered to a nursing
	woman.
STOT-single Exposure	Systemic absorption of this product may result in toxic effects on the CNS and the cardiovascular system.
	Adverse effects could be more pronounced in those individuals with pre-existing diseases of central nervous system or cardiovascular system or those receiving medications that affect these systems (such as antihypertensive agents, antiarrhythmic agents or CNS depressant medications). Effects could also be more pronounced in individuals with a compromised ability to metabolize and clear active ingredients from the blood and body tissues (such as severe liver or kidney disease).
	Prilocaine may cause methemoglobinemia in high doses and so may aggravate congenital or idiopathic methemoglobinemia.
STOT-repeated Exposure	Chronic effects are unlikely to occur. Repeated exposure to high levels of an amide anesthetic in animals produced adverse effects on the liver and CNS.
	Prilocaine may cause methemoglobinemia in high doses and so may aggravate congenital or idiopathic methemoglobinemia.
Aspiration Hazard	Aspiration hazard is low. May cause tingling/numbness in exposed areas (paresthesia). Intratracheal (rabbit) LD50 for prilocaine is 65 mg/kg

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o-toluidine is a carcinogen in both species. The lowest dose corresponds to approximately 50 times the maximum amount of *o*-toluidine to which a 50 kg subject would be expected to be exposed following a single injection (8 mg/kg) of prilocaine.

Studies in rats with 2,6-xylidine indicated carcinogenic potential of this metabolite of lidocaine at high doses.

Reproductive Toxicity:

Prilocaine: Reproduction studies have been performed in rats at doses up to 30 times the human dose and revealed no evidence of impaired fertility or harm to the fetus.

Lidocaine: No teratogenic effects were noted in embryo-fetal development studies in which rats or rabbits were treated during the period of organogenesis. Embryotoxicity was seen in rabbits, at maternally toxic doses. In rats, decreased pup survival was seen for dams treated during late pregnancy and lactation, at a dose that was maternally toxic and affected the duration of gestation.

Lidocaine and prilocaine: No effects on embryofetal development were seen in a study in which lidocaine and prilocaine were given in combination, during organogenesis.

There are, however, no adequate and well-controlled studies in pregnant women. Animal reproduction studies are not always predictive of human response. General consideration should be given to this fact before administering prilocaine to women of childbearing potential, especially during early pregnancy when maximum organogenesis takes place.

Lidocaine, and in all probability, prilocaine are excreted in breast milk in small amounts. However it is unlikely that effects will be seen in the child following treatment with Oraqix. Thus breast-feeding can be continued following treatment, with

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	Intraperitoneal (mouse) LD50: 30 mg/kg	
	Subcutaneous (mouse) LD50: 632 mg/kg	
	Intravenous (mouse) LD50: 55 mg/kg	
	Intravenous (guinea pig) LD50: 20 mg/kg	
	Intravenous (rabbit) LD50: 18 mg/kg	
	Intratracheal (rabbit) LD50: 65 mg/kg	
	Altered sleep-time, convulsions recorded.	
	Tittered sleep time, convensions recorded.	
	Only selected data are presented here. See actual	
	entry in RTECS for complete information.	
Skin Corrosion / Irritation:		
	May cause mild skin irritation.	
Serious Eye Damage / Irritation	May cause irritation, excessive watering	
	(lacrimation) and eye damage, blurred vision and	
	numbness.	
Respiratory or Skin	Repeated or prolonged contact may cause	
Sensitisation:	sensitization in a small proportion of the	
	population. May cause numbness.	
Germ Cell Mutagenicity:	Studies of prilocaine in animals to evaluate the	
	mutagenic potential have not been conducted.	
	O-toluidine (0.5 mg/mL), a metabolite of	
	prilocaine, showed positive results in Escherichia	
	coli DNA repair and phage-induction assays. Urine	
	concentrates from rats treated with <i>o</i> -toluidine (300)	
	mg/kg, orally) were mutagenic for Salmonella	
	typhimurium with metabolic activation. Several	
	other tests, including reverse mutations in five	
	different Salmonella typhimurium strains with or	
	without metabolic activation and single strand	
	breaks in DNA of V79 Chinese hamster cells, were	
	negative.	
	Genotoxicity tests with lidocaine were negative.	
	However, whilst Ames genotoxicity tests with	
	2,6-xylidine were negative a chromosome	
	aberration test in CHO cells indicated an in vitro	
	genotoxic potential of this metabolite of lidocaine.	
Carcinogenicity:	Studies of prilocaine or lidocaine in animals to	
	evaluate the carcinogenic potential have not been	
	conducted.	
	Chronic oral toxicity studies of <i>o</i> -toluidine, a	
	metabolite of prilocaine, in mice (150–4800	
	mg/kg) and rats (150–800 mg/kg) have shown that	
	mg/kg/ and rais (130-000 mg/kg/ nave shown that	

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11	TOXICOL	OCICAL	INFORMA	TION
11.	IVAICUL			

Acute Toxicity:

LD50 / LC50 Mixture: Unknown

Lidocaine hydrochloride (Readily available toxicity data unavailable for base form):

Intravenous / child

Lowest published toxic dose: 60 mg/kg/1 hour Behavioral: Convulsions or effect on seizure

threshold

Vascular: BP lowering not characterized in

autonomic section

Intravenous / infant

Lowest published toxic dose: 10 mg/kg Behavioral: Convulsions or effect on seizure

threshold. Coma

Lung, Thorax, or Respiration: Other changes

Intravenous / man

Lowest published toxic dose: 9 mg/kg/4 hour-

continuous

Cardiac: Cardiomyopathy including infarction

Intravenous / man

Lowest published toxic dose: 7.143 µg/kg Cardiac: Pulse rate increased without fall in BP

Oral / infant

Lowest published toxic dose: 1.632 mg/kg/1 week-

intermittent

Behavioral: Somnolence (general depressed activity). Convulsions or effect on seizure

threshold

Prilocaine hydrochloride (Readily available toxicity data unavailable for base form):

Parenteral (man) LDLo: 12.43 mg/kg/1h - I Nil

Reported

Intraperitoneal (rat) LD50: 148 mg/kg Subcutaneous (rat) LD50: 790 mg/kg Intravenous (rat) LD50: 56.6 mg/kg

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9. PHYSICAL AND CHEMICAL PROPERTIES		
Appearance:	Clear aqueous solution	
Odour (odour threshold):	Odorless	
pН	3.3 - 5.5	
Melting Point:	Not available	
Initial Boiling Point:	Not available	
Boiling Range:	Not available	
Flash Point:	Non Combustible	
Evaporation Rate:	Not available	
Flammability:	Not flammable	
Upper and Lower Flammability	Not Relevant	
Limits		
Vapour Pressure:	Not available	
Vapour Density:	Not available	
Relative Density:	1.0	
Solubility(ies):	Not applicable	
Partition Coefficient (n-octanol	Not available	
/ water):		
Auto-ignition Temperature:	Not applicable	
Decomposition Temperature:	Not available	
Viscosity:	Not available	

10. STABILITY AND REACTIVITY		
Reactivity:	Non-reactive	
Chemical Stability:	Product is considered stable under normal	
	conditions	
Possibility of Hazardous	Unlikely unless in contact with alkaline conditions	
Reactions:		
Conditions to Avoid:	Open burning/incineration	
Incompatible Materials:	Compounds that react violently with water. Strong	
	reducing agents.	
Hazardous Decomposition	Fumes of Carbon Monoxide, Carbon Dioxide,	
Products:	Nitrogen Oxides Hydrogen Chloride gas	

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7. HANDLING AND STORAGE		
Handling:	No special precautions are necessary when	
	handling packed product. In case of release, avoid	
	contact with skin and eyes. Do not breathe mist.	
Storage:	Protect from light. Store in original containers and	
	packaging as recommended by manufacturer. Keep	
	containers securely sealed and cool. Store below	
	25°C. Check that containers are clearly labelled.	

8 EXPOSURE CONTROL	LS / PERSONAL PROTECTION
Exposure Control Limits:	No exposure limits assigned for product. Prilocaine hydrochloride - 5 mg/m³ COM, REL TWA
Special Protective Measures:	Wear suitable protective clothing
	Eye: Chemical goggles or face shield.
	Hands/feet: Wear chemical protective gloves, e.g. PVC.
	Other: Laboratory coat and P.V.C. apron.
	Engineering controls: Use in a well-ventilated area. General exhaust is adequate under normal operating conditions. Local exhaust ventilation may be required in specific circumstances. If needed, use a NIOSH approved respirator for vapors, dusts and mists with TLV greater than 0.05 mg/m ³ .
	Respiratory Protection: Material does not require special ventilators, respirators, etc.
	Work Hygienic Practices: Avoid ingestion & contact with eyes. Remove / launder contaminated clothing & shoes before reuse. Wash hands after use.
	Supplemental Health & Safety Information: Irritating to the eye. Contact may also cause numbness and loss of sensation.

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4. FIRST AID MEASURES			
Eye Contact:	Flush immediately with eye wash solution or clean		
	water, holding the eyelids apart, for at least 15		
	minutes. Obtain medical attention.		
Skin Contact:	Remove contaminated clothing. Wash skin with		
	soap and water. If symptoms (irritation or		
	blistering) occur obtain medical attention		
Inhalation:	Remove patient from exposure. Obtain medical		
	attention if ill effects occur. May cause		
	tingling/numbness in exposed areas (paresthesia).		
	High atmospheric concentrations may lead to		
	anaesthetic effects.		
Ingestion:	Do not induce vomiting. Rinse mouth with water		
	and give 200-300 ml of water to drink		
	(8-10 ounces). Never give anything by mouth if		
	unconscious. Obtain medical attention.		
	May produce numbness of the tongue and		
	anesthetic effects on the stomach. Ingestion of 5 to		
	25 mL of 2% viscous Xylocaine (lidocaine) has		
	resulted in seizures in children.		

5. FIRE FIGHTING MEASURES				
Suitable Extinguishing Media:	Use appropriate agent for involved fire (i.e., water spray, carbon dioxide, dry chemical powder or appropriate foam).			
Specific Hazards Arising from	If involved in a fire, it may burn and emit noxious			
the Chemical(s):	and toxic fumes.			
Protection of Fire-fighters:	A self contained breathing apparatus and suitable protective clothing should be worn in fire			
	conditions.			

6. ACCIDENTAL RELEASE MEASURES				
Personal Precautions:	Ensure suitable personal protection during removal of spillages. Take care to avoid needles and broken			
	containers. Clean spills with normal procedures used for non-hazardous liquids.			
Environmental Precautions:	Transfer spilled vials to a suitable container for disposal. Sweep/soak up, place in a bag and hold for waste disposal.			
Containment and Clean up:	Clear up spillages. Wash the spillage area with water. Transfer spilled vials to a suitable container for disposal. Ventilate area and wash spill site after material pickup is complete.			

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1. IDENTIFICATION (MATERIAL AND MANUFACTURER)					
Product Name:	Oraqix®				
Synonym(s):	Lidocaine and Prilocaine periodontal gel				
Product Use:	Indicated for adults who require localized anesthesia in				
	periodontal pockets during scaling and/or root planing.				
Manufacturer / Supplier:	DENTSPLY Pharmaceutical				
	1301 Smile Way				
	York, PA 17404				
	USA				
	Telephone number: 1-800-225-2787				
	Fax number: 717-699-4148				
Emergency telephone numbers:					
Country		Call	Phone Number		
		Order			
USA		Primary	717-767-8523		
			717-887-9723		
		Secondary:	717-767-4120		
			717-495-5901		
Canada		Primary	1-800-263-1437		

2. HAZARD IDENTIFICATION				
Hazard Classification:	Xn; R22 Carc3; R40 R43			
GHS Hazard Labelling:				
Canadian Hazard Warning:				
Other Hazards:	None			

3. COMPOSITION / INFORMATION ON INGREDIENTS				
Name	CAS Number	% conc		
Lidocaine base	137-58-6	2.5		
Prilocaine base	721-50-6	2.5		

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