SUMMARY OF PRODUCT CHARACTERISTICS

NAME OF THE MEDICINAL PRODUCT

Package leaflet: Information for the user Orabloc®1:200,000
Articaine hydrochloride and adrenaline (epinephrine) solution for injection 1:200,000

Read all of this leaflet carefully before this medicine is given to you by your dentist because it contains important information foryou.

Keep this leaflet. You may need to read it again.
 If you have any further questions, ask your dentist, doctor or pharmacist.

If you get any side effects, talk to your doctor, dentist or pharmacist. This includes any possible side effects not listed in this leaflet.

What is in this leaflet:

1. What Orabloc is and what it is used for

2. What you need to know before you are given Orabloc

3. How Orabloc is used

Possible side effects 5. How to store Orabloo

Contents of the pack and other information

 Non-tents of the pack and other miorination.

1. What Orabloc is and what it is used for:

Orabloc is a medicine for local anaesthetization (local anesthetic) in dentistry.

It contains the active ingredients articaine (local anesthetic) and epinephrine (adrenaline).

Epinephrine (adrenaline) narrows the blood vessels. This reduces blood flow at the point at which you receive the needle from vour doctor(local anemia).

This reduces bleeding in treatments and extends the effect of the local anaesthetization

Orabloc is used for the local anaesthetisation of dental treatments in adults, adolescents and children from 4 year old, including: -uncomplicated removal of one or several teeth,

- uncomplicated reintoval of one of several teetin,
- removal of caries,
- crown stump preparations.

2. What you need to know before you are given Orabloc

- if you are allergic to articaine, or other local anaesthetic of the acid amide type, to epinephrine (adrenaline) or any of the other

ingredients of this medicine (listed in section 6), if you suffer from severe heart rhythm disorders (e.g. second and third-degree AV block),

I fyou have a very low pulse, if you have a very low pulse, if you have a very low pulse, if you suffer from acute heart failure (acute heart weakness, e.g. after myocardial infarction (e.g. heart attack)), if you have very low blood pressure, if you have very low blood pressure, if you are a bronchial asthmatic and suffer from sulphite hypersensitivity (asthma attacks, triggered by sulphite).

- in children under 4 years of age
Owing to the effects of the epinephrin (adrenaline) fraction. Orabloc must not be used on you:

if local anaesthetisation in a 'tenninal area' of the blood vessels is being carried out (i.e. in an area supplied with blood only by - in local arraesureusation in a terminar area of the the branches of a single artery),
- if you have raised intraocular pressure (glaucoma),
- if you have an overactive thyroid,

if you suffer from sudden-onset tachycardia (paroxysmal tachycardia),

if you suffer from a particular form of heart rhythm disorders (absolute arrhythmia with high pulse rate), if you had a myocardial infarction (e.g. heart attack) in the last 3 to 6 months,

If you had coronary artery bypass surgery in the last 3 months,

if you take certain beta blockers, such as propranoiol. There is the danger of a hypertensive crisis (very high blood pressure) or severe slowing of the pulse,

if you suffer from phaeochromocytoma (adrenaline-producing tumour generally in the adrenal medulla),

-if you suffer from phaseochromocyforma (adrenaline-producing turnour generally in the adrenal medulla),
 -if you have very high blood pressure,
 -if you have very high blood pressure,
 -if you are treated simultaneously with certain medicines for the treatment of depression and Parkinson's disease (tricyclic antidepressants, MAO inhibitors). These medicines may intensify the cardiovascular effects of epinephrine (adrenaline). This may apply up to 14 days after the conclusion of treatment with MAO inhibitors.
 Orabloc must not be administered intravenously (into a vein).
 Warnings and precautions
 Talk to your dentist or pharmacist before Orabloc is used:
 -if you have a deficiency of a specific enzyme (cholinesterase deficiency). This may lead to a slowed and possibly intensified action of Orabloc.

- if there is local inflammation or infection at the site at which the syringe is to be inserted. Increased absorption of Orabloc takes place in this event, with efficacy being reduced

suffer from epilepsy (see section 4).

pract in inservent, with embacty being reduced.

Orabloc may be administered only after careful medical examination if you:

- suffer from disturbances of blood clotting,

- suffer from severe kidney or liver insufficiency (e.g. in the case of nephritis or cirrhosis of the liver),

- are receiving concomitant treatment with halogenated inhalation anaesthetics (see Using other medicines),

If you suffer from any of the following diseases, Orabloc may be administered only after careful medical examination:

cardiovascular diseases, e.g.:
angina pectoris (disturbance of blood flow to the heart with pain and feeling of tightness in the chest),

- arigina pecions (culturalized or Joulou low to ure near with pain and reening or agrintess in the chest),
- arteriosclerosis (narrowing of arteries by deposits, e.g. by blood fats),
- heart failure (heart muscle weakness),
- coronary heart disease (narrowing of the coronary vessels),
- angina pectoris (disturbance of blood flow to the heart with pain and feeling of tightness in the chest),

after a myocardial infarction (eq heart attack). heart rhythm disorders (irregular pulse).

high blood pressure

ders of blood flow to the brain

- disorders of blood now of the brain, - after a stroke, - chronic bronchitis, pulmonary emphysema (pathological overinflation of the lungs), - diabetes (diabetes mellitus),

severe anxiety states.

To avoid side effects, your dentist will

check your medical history and concomitant treatments

check your medical history and concomitant treatments,
 practise a test injection if there is a risk of allergy to the product,
 choose the dosage as low as possible,
 before the injection, carefully check that he has not struck a blood vessel.
 Use of the product can result in protonged numbness in the area of the mouth after the dental procedure: in young children, care should be taken to avoid self-biting, which could result in soft tissue injury.

Other medicines and Orabloc Tell your dentist or pharmacist if you are taking, have recently taken or might take any other medicines

If you have simultaneously used other medicines for local anaesthetisation, their effects on the cardiovascular system and nervous system may be intensified.

Orabloc contains the active ingredient epinephrine (adrenaline). This active ingredient narrows the blood vessels and raises blood pressure. The blood pressure-raising effect of epinephrine (adrenaline) may be intensified by certain medicines for the treatment of depression and Parkinson's disease (e.g. tricyclic antidepressants and MAO inhibitors must therefore not be taken at the same time). (Please note section «2. What you need to know before you use Orabloc»).

Please also note the section «Do not use Orabloc» concerning the simultaneous use of certain beta blockers (such as propranolol). Epinephrine (adrenaline) may inhibit the release of insulin from the pancreas. This may reduce the effect of oral antidiabetics (medicines for the treatment of diabetes).

(medicines for the treatment of diabetes).

If Orablo is used at the same time with certain anaesthetic gases (e.g. halothane), this may trigger heart rhythm disturbances. Phenothiazines can influence the blood-pressure-increasing effects of epinephrine (adrenaline). Therefore concomitant treatment should be avoided. If concomitant treatment is necessary patients should be monitored carefully. Please note: In patients treated with blood cloi-inhibiting (blood-thinnings) medicines (such as heparin or acetylsalicylic acid), an inadvertent injection into a blood vesselas part of local anaesthesia may lead to serious bleeding and also the tendency to bleed

(danger of bleeding) may also be generally increased.

(danger of bleeding) may also be generally increased.

Orabloe with food and drink

After using Orabloe, you should not consume any food until the local anaesthetic has worn off.

Pregnancy and breast-feeding
If you are pregnant, your dentist should use Orabloc only after a careful risk-benefit assessment.

For articeine, there is no experience of use during pregnancy, other than at birth. Animal experimental studies have shown that epinephrine (adrenaline) has a harmful effect on progeny at doses higher than used for dental anesthesia. After administration by mistake of Orabloc to a blood vessel in the mother, a reduction in blood flow to the uterus may occur as a result of the epinephrine

(additionally fraction).

The active ingredients of Orabloc are broken down quickly by your body. This means that quantities of the active ingredients that are harmful to the breastfed infant are not excreted into breast milk. You need not therefore intem1pt breastfeeding in connection with short-term treatment with Orabloc.

Your dentist will decide when you may drive or use machines again after an intervention. In relevant investigations, no impairment of normal ability to drive was identified after local anaesthetisation with articaine.

Orabloc contains sodium metabisulphite (E223) and sodium.
This medicinal product contains less than 1 mmol sodium (23 mg) per dose, i.e. essentially sodium-free.

3. How Orabloc is used

Your dentist determines the dosage and method of administration of Orabloc. He will generally be guided by the following Dosage
An injection of 1.8 ml Orabloc per tooth is generally enough for the simple removal of uninflamed upper jaw teeth. Ina few cases, a subsequent injection of 1 - 1.8 ml may be necessary to achieve full local anaesthetisation. Generally, not every tooth needs to be anaesthetised with the full quantity of Orabloc for the removal of neigh bouring teeth. The

number of injections can normally be reduced. If an incision or a suture on your palate is necessary, an injection of approx. 0.1 ml per puncture is sufficient. When removing uninflamed front molars in the lower jaw, an injection of 1.8 ml per tooth is generally sufficient. If a full effect does not arise after that, your dentist can give a subsequent injection of 1 - 1.8 ml. Only if the site is not completely anaesthetised even after the subsequent injection your dentist can carry out the otherwise susual anaesthetisation of the entire mandibular nerve.

Depending on the scale and length of the treatment, 0.5 - 1.8 ml Orabloc 1:200,000 solution for injection is sufficient for the removal

of caries in preparation for a filling and for crown polishing.

This does not apply, however, to lower-law molars.

Inis does not apply, however, to lower-jaw molars.

Adults can receive up to 7 mg articaine per kg body weight in the course of a treatment. Quantities of up to 500 mg (equivalent to 12.5 ml solution for injection) are normally tolerated well.

Elderly and patients with severe disturbance of liver and kidney function In elderly patients and patients with severe disturbance of liver and kidney function (e.g. in the case of nephritis or cirrhosis of the liver), increased quantities of articaine mayarise in the blood. If you are in these patient groups, your dentist should take particular care to ensure that the smallest possible quantity for adequate anaesthetisation is used.

If Orabloc is used in children and adolescents, the minimum volume necessary to achieve adequate anaesthesia should be used. If Orabloc is used in children and adolescents, the minimum volume necessary to achieve adequate anaesthesia should be used. The injection quantity is to be dosed individually according to the child's and adolescent's age and weight. A maximum dose of 5 mg articaine per kg body weight should not be exceeded.

This product has not been studied in children less than I year old. Method of administration

Orabloc is intended for administration in the oral cavity (dental use).

To prevent infections (e.g. transmission of hepatitis), new and sterile syringes and needles must be used for each injection. For single use. Any unused solution should be discarded.

If desolvered resolved the application and the used.

f discoloured or cloudy, the medicine must not be used

If you received too much of Orabloc

essive quantities of Orabloc are used, disturbances of nervous system function may occur, e.g.:

seizures and disorders of breathing.

- seizures and disorders of breathing.

Disturbances of cardiovascular function may also arise, such as a fall or rise in your blood pressure. Such disturbances require dental monitoring and possibly suitable therapy by your dentist.

If you have any further questions on the use of this medicine, ask your dentist, doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them. Common (may affect up to 1 in 10

people)
Nausea, vomiting, disturbed sensation of touch (paraesthesia), reduction in sensitivity of sensation in the mouth and facial area (hypoaesthesia); headaches which are presumably attributable to the epinephrine fraction Uncommon (may affect up to 1 in 100

Repid heart beat, (lachycardia), dizziness
Unknown (frequency cannot be estimated from the available data)

- Heart rhythm disturbances, increase in blood pressure, drop in blood pressure, lowered heart rate (bradycardia), heart failure and

- Depending on dose (particularly in the case of excessive dosing or inadvertent injection into ablood vessel), disturbances of the

nervous system may arise, e.g.:
- agilation, nervousness,
- drowsiness ranging to loss of consciousness, coma,
- respiratory disturbances ranging to respiratory arrest,
- muscle tremor, muscle twitching ranging to convulsions.
- During or shortly after the injection of local anaesthetics in the head area, temporary visual disorders (blurred vision, blindness, - Nerve damage (e.g. of the facial nerve) and a reduction in sensitivity of taste in the mouth and facial area are not side effects that

- Hypersensitivity reactions (allergic or allergy-like reactions) may arise. These may manifest themselves as swelling or inflammation at the injection site. Hypersensitivity reactions arise that are not restricted to the injection site:

- continective ussue minimitudin,
- common cold,
- facial swelling (in the form of Quincke's oedema) with swelling of the upper and/or lower lip and/or cheeks,
- swelling of the laryngeal area with a feeling of tightness and swallowing complaints,

respiratory complaints ranging to anaphylactic shock. -respiratory complaints ranging to anaphylactic shock.
- Owing to the content of sodium metabisulphite hypersensitivity reactions that may manifest themselves as vomiting, diarrhoea, wheezing, acute asthma attacks, disturbances of consciousness or shock may arise very rarely, particularly in bronchial asthmatics.
-respiratory dystunction (tachypnea, bradypnea) that may lead to apnea
Additional side effects in children
In young children, compared to adults, there is an increased risk of self-bitting, which could result in soft tissue injury, because of prolonged numbness in the area of the mouth after the dental procedure.
If a side effect arises suddenly or develops strongly, inform a doctor immediately. This is particularly important because certain pharmaceutical side effects (e.g. fall in blood pressure or respiratory disturbances) may become life-threatening.
Reporting of side effects

pharmaceutical side effects (e.g. fall in blood pressure or respiratory disturbances) may become life-threatening.

Reporting of side effects

If you get any side effects, talk to your doctor, dentist or pharmacist. This includes any possible side effects not listed in this leaflet.

By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Orabloc

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton and on the cartridge after «EXP». The expiry date refers to the last day of that month

to the last day of that mount.

Do not store above 25°C. Store in the original package in order to protect from light.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help to protect the environment.

6. Contents of the pack and other information

The active substances are articaine hydrochloride and adrenaline (epinephrine).

- 1 ml solution for injection contains 40 mg of articaine hydrochloride and 0.005 mg adrenaline (epinephrine) as adrenaline tartrate.

- One cartridge of 1.8 ml solution for injection contains 72 mg articaine hydrochloride and 0.009 mg of adrenaline (epinephrine) as

One cartridge of 1.8 ml solution for injection contains 72 mg articaine hydrochloride and 0.009 mg of adrenaline (epinephrine) as adrenaline latrate.
 The other ingredients are: sodium metabisulphite (E223), sodium chloride, hydrochloric acid 2% (for pH adjustment), water for injection.
 What Orabloc looks like and contents of the pack
 Orabloc is a sterile, clear, colourless solution for injection in clear glass cartridges closed at one end with a bromobutylic rubber

plunger and at the other with an aluminium cap and rubber seal The cartridges are packaged in blisters (10 cartridges/blister); the blisters are packaged in a cardboard box containing 5 x 10 or 10 x 10 cartridges.

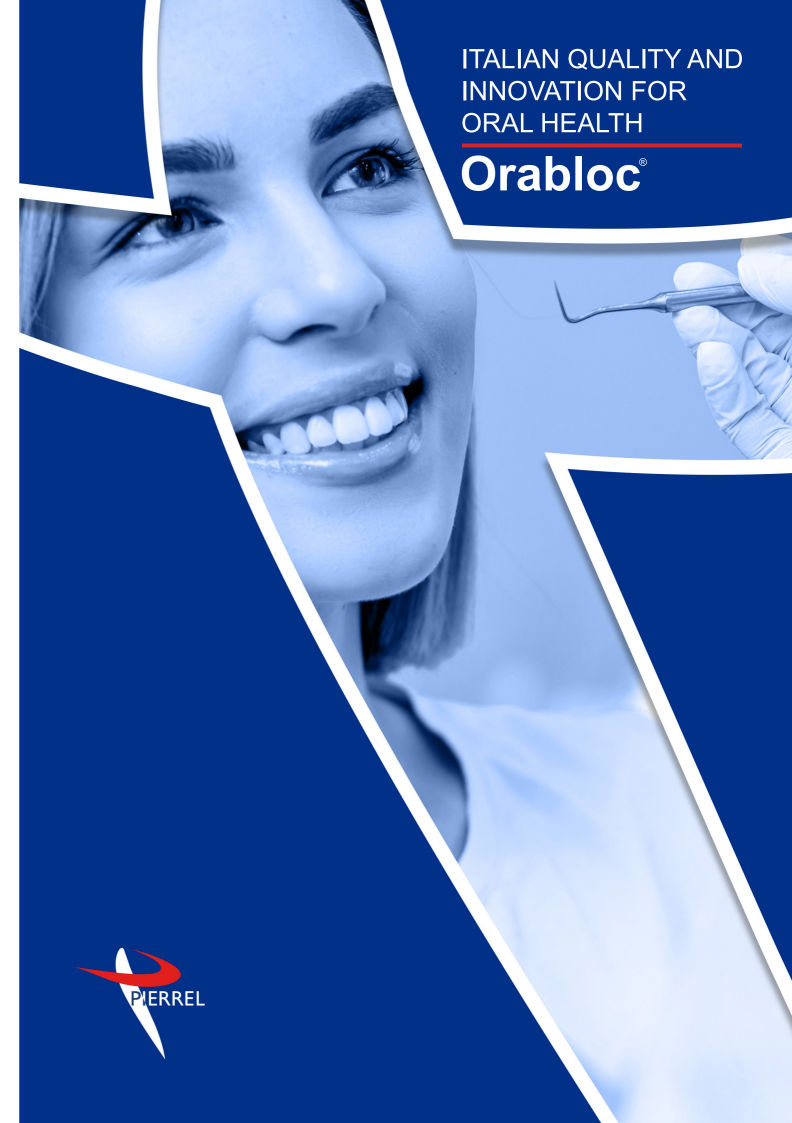
10 x 10 cartridges. Each cartridge is assembled in a plastic injector; each injector containing a cartridge is placed in a sealed blister; the injectors are packaged in a cardboard box together with an Instructions for use of the injector: 50 or 100 units per commercial pack. Not all pack sizes may be marketed.

da Statale Appia, 7 BIS 46/48 – 81043 Capua (CE)

This medicinal product is authorised in the Member States of the EEA under the following names: Germany Articainhydrochlorid mit Epinephrin Pierrel 40 mg/ml + 0,005 mg/ml] Injektionslösung Austria Orabloc 40 mg/ml + 5 Mikrogramm/ml Injektionslösung France Orabloc 40 mg/ml Adrenalinee au 1/200 000 Solution injectable

his leaflet was last revised in November 2024

Marketing Authorisation Holder and Manufacturer



Orabloc[®]

Manufactured in Italy by: Pierrel S.p.A. Strada Statale Appia 46/48 - 81043

Articaine HCI 4% and epinephrine 1:100,000 and epinephrine 1:200,000. Injection.

- » Rapid onset of anesthesia within 1-3 minutes.
- » Complete anesthesia lasts about 1 hour for infiltrations, up to 2 hours for nerve block.
- 10% overage of epinephrine¹
- 24 month shelf life at room temperature.
- Sodium edetate free, methylparaben free and latex free.
- » Most common adverse reactions (incidence >2%) are headache and pain.
- » Each cartridge is sealed individually in the blister for maximum protection.



Orabloc is indicated for local, infiltrative, or conductive anesthesia in both simple and complex dental procedures:

- » For most routine dental procedures, Orabloc containing epinephrine 1:200.000 is preferred.
- » When more pronounced homeostasis or improved visualization of the surgical field are required, Orabloc containing epinephrine 1:100,000 may be used.

Orabloc have a 24 month shelf life

- » Store at room temperature; 25°C (77°F), with brief excursions permitted between 15°C (59°F) and 30°C(86°F).
- Protect from light.
- Do not freeze.

Orabloc packaging

- » Each cartridge is individually sealed for maximum protection up to the moment of use.
- Cartridges packed 10 to a blister tray to avoid glass to glass contact
- » Blister trays packaged in boxes of 50.

Dosage and administration – Adults

- » For normal healthy adults, the maximum dose of Orabloc administered by submucosal infiltration and/or nerve block should not exceed 7mg/kg (0.175 mL/kg) of articaine HCl.
- » Dosage should be reduced in elderly patients and in patients with cardiac or liver disease.

Pediatric patients ages 4 to 16 years

- » The quantity of Orabloc in children ages 4 to 16 years of age to be injected should be determined by the age and weight of the child and the magnitude of the operation.
- The maximum dose of Orabloc should not exceed 7 mg/ kg (0.175 mL/kg) of articaine HCI (see Use in Specific Populations). Use in pediatric patients under 4 years of age is not recommended.

The American Heart Association (AHA) recommends using the lowest possible quantity of eninephrine (Kaplan EL ed. Cardiovascular disease in dental practice. Dallas, TX: American Heart Association, 1986

IMPORTANT SAFETY INFORMATION

Care should be taken to avoid accidental intravascular injection, which may be associated with convulsions followed by coma and respiratory arrest. Local anesthetic solutions that contain a vasoconstrictor should be used cautiously, especially in patients with impaired cardiovascular function or vascular disease. Administration of Orabloc results in a 3 to 5 fold increase in plasma epinephrine concentrations compared to baseline. However, in healthy adults it does not appear to be associated with marked increases in blood pressure or heart rate, except in the case of accidental intravascular injection. The most common adverse reactions (incidence >2%) are headache and pain. Inform patients in advance of the possibility of temporary loss of sensation and muscle function following infiltration and nerve block injections. Instruct patients not to eat or drink until normal sensation returns.

Please see accompanying full prescribing information or visit www.orabloc.com

Orabloc is an amide local anesthetic containing a vasoconstrictor indicated for local, infiltrative, or conductive anesthesia in both simple and complex dental procedures. Orabloc contains sodium metabisulfite.

Orabloc is contraindicated in patients who are hypersensitive to products containing sulfites. Products containing sulfites may cause allergic-type reaction including anaphylactic symptoms and life-threatening or less severe asthmatic episodes in certain susceptible people. Please to see or download the full prescribing information visit www.orabloc.com



Please see accompanying full prescribing information or visit www.orabloc.com using the QR code.

To learn more about Orabloc call Pierrel at 1-610-989-4222 or email to orabloc@pierrelgroup.com

SUMMARY OF PRODUCT CHARACTERISTICS

NAME OF THE MEDICINAL PRODUCT

Orabloc 1:100,000
Articaine hydrochloride 4% with adrenaline (epinephrine) solution for injection 1:100,000

Read all of this leaflet carefully before this medicine is given to you by your dentist because it contains important information

tur you.

Keep this leaflet. You may need to read it again.

If you have any further questions, ask your dentist, doctor or pharmacist.

If you get any side effects, talk to your doctor, dentist or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

- What is in this leaflet:

 1. What Orabloc is and what it is used for 2. What you need to know before you are given Orabloc
- 3. How Órabloc is used 4 Possible side effects
- 5. How to store Orabloo
- 6. Contents of the pack and other information

 1. What Orabloc is and what it is used for:
- Orabloc is a medicine for local anaesthetization (local anesthetic) in dentistry.
- It contains the active ingredients articaine (local anaesthetic) and epinephrine (adrenaline). Epinephrine (adrenaline) narrows

the blood vessels. This reduces blood flow at the point at which you receive the needle from your doctor (local anemia). This reduces bleeding in treatments and extends the effect of the local anaesthetization.

Totablo is used for the local anaesthetization of dental treatments in adults, adolescents and children from 4 year old, including:

- surgery of the oral mucosa or bone in which a greater reduction in blood flow is important,

- surgery of the dental pulp,

- longer surgical interventions
- oral surgery of the bone with open gums.
- moval of cysts (fluid-filled cavities in the tissue) surgery in the gum area or at the gum-tooth margin

- 2. What you need to know before you are given Orabloc Do not use Orabloc
 -if you are allergic to articaine, or other local anesthetic of the acid amide type, to epinephrine (adrenaline) or any of the other
- ingredients of this medicine (listed in section 6), if you suffer from severe heart rhythm disorders (e.g. second and third-degree AV block),
- if you suffer from acute heart failure (acute heart weakness, e.g. after myocardial infarction (e.g. heart attack)),

- if you suffer from acute heart failure (acute heart weakness, e.g. after myocardial infarction (e.g. heart attack)),
- if you have very low blood pressure,
- if you have very low blood pressure,
- if you are a bronchial asthmatic and suffer from sulphite hypersensitivity (asthma attacks, triggered by sulphite).
- in children under 4 years of age
- Owing to the effects of the epinephrin (adrenaline) fraction, Orabloc must not be used on you:
- if local anaesthetisation in a 'tenninal area' of the blood vessels is being carried out (i.e. in an area supplied with blood only

- by the branches of a single artery). ular pressure (glaucoma)

- in you have nased initiational pressure (glaucoma), if you have an overactive thyrrioid, if you suffer from sudden-onset tachycardia (paroxysmal tachycardia), if you suffer from a particular form of heart rhythm disorders (absolute arrhythmia with high pulse rate), if you had a myocardial infarction (e.g. heart attack) in the last 3 to 6 months,
- if you had coronary artery bypass surgery in the last 3 months
- if you take certain beta blockers, such as propranolol. There is the danger of a hypertensive crisis (very high blood pressure)
- or severe slowing of the pulse.
- if you suffer from phaeochromocytoma (adrenaline-producing tumour generally in the adrenal medulla)
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 if you have very high blood pressure,
 if you are treated simultaneously with certain medicines for the treatment of depression and Parkinson's disease (tricyclic antidepressants, MAO inhibitors). These medicines may intensify the cardiovascular effects of epinephrine (adrenaline). This may apply up to 14 days after the conclusion of treatment with MAO inhibitors.

Orabloc must not be administered intravenously (into a vein).

Warnings and precautions

- varinings and precadurons

 Talk to your dentist or phannacist before Orabloc is used:

 if you have a deficiency of a specific enzyme (cholinesterase deficiency). This may lead to a slowed and possibly intensified action of Orabloc

 if there is local inflammation or infection at the site at which the syringe is to be inserted. Increased absorption of Orabloc takes
- place in this event, with efficacy being reduced.
- Orabloc may be administered only after careful medical examination if you:

- Orabloc may be administered only after careful medical examination if you:

 suffer from disturbances of blood clotting,
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 are receiving concomitant treatment with halogenated inhalation anaesthetics (see Using other medicines),
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 If you suffer from any of the following diseases, Orabloc may be administered only after careful medical examination and your denists should consider using Orabloc 1200,000

 Articaine hydrochloride 4% with adrenaline (epinephrine) solution for injection 1200,000 instead of Orabloc 1:100,000 Articaine hydrochloride 4% with adrenaline (epinephrine) solution for injection 1:100,000 as the former contains less epinephrine
- adrename).
 cardiovascular diseases, e.g.:
 angina pectoris (disturbance of blood flow to the heart with pain and feeling of tightness in the chest),
- arteriosclerosis (narrowing of arteries by deposits, e.g. by blood fats) heart failure (heart muscle weakness),
- coronary heart disease (narrowing of the coronary vessels) anging pectoris (disturbance of blood flow to the heart with pain and feeling of tightness in the chest).

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- chronic bronchitis, pulmonary emphysema (pathological overinflation of the lungs). diabetes (diabetes mellitus)

- clauderes (valueus minimus),
 severe anxiety states.
 To avoid side effects, your dentist will
 check your medical history and concomitant treatments,
 practise a test injection if there is a risk of allergy to the product,

- choose the dosage as low as possible,
 before the injection, carefully check that he has not struck a blood vessel.
 Use of the product can result in prolonged numbness in the area of the mouth after the dental procedure: in young children, care should be taken to avoid self-biting, whichcould result in soft tissue injury.

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Other medicines and Orabloc

Tell your dentist or phannacist if you are taking, have recently taken or might take any other medicines.

If you have simultaneously used other medicines for local anaesthetisation, their effects on the cardiovascular system and nervous system may be intensified.

Orablic contains the active ingredient epinephrine (adrenaline). This active ingredient narrows the blood vessels and raises blood pressure. The blood pressure-raising effect of epinephrine (adrenaline) may be intensified by certain medicines for the treatment of depression and Parkinson's disease (e.g. tricyclic antidepressants and MAO inhibitors must therefore not be taken at the same time). (Please note sections.2. What you need to know before you use Orabloca).

Please also note the section «Do not use Orabloc» concerning the simultaneous use of certain beta blockers (such as propranolol). Epinephrine (adrenaline) may hibit the release of insulin from the pancreas. This may reduce the effect of oral antidiabetics (medicines for the treatment of diabetes). If Orabloc is used at the same time with certain anaesthetic gases (e.g. halothane), this may trigger heart rhythm disturbances. Phenothizainse can influence the blood-pressure-increasing effects of epinephrine (adrenaline). Therefore concomitant Phenothiazines can influence the blood-pressure-increasing effects of epinephrine (adrenaline). Therefore concomitant

renormazines can imiuence the blood-pressure-increasing effects or epineprinne (acrealine). Therefore concomitant treatment should be avoided. If concomitant treatment is necessary patients should be monitored carefully. Please note: In patients treated with blood clot-inhibiting (*blood-thinning*) medicines (such as heparin or acetylsalicylic acid), an inadvertent injection intoa blood vessel as part of local anaesthesia may lead to serious bleeding and also the tendency to bleed (danger of bleeding) may also be generally increased.

Orabloc with food and drink

Orabloc with food and drink
After using Orabloc, you should not consume any food until the local anaesthetic has worn off.

Pregnancy and breast-feeding
If you are pregnant, your dentist should use Orabloc only after a careful risk-benefit assessment.

For articaine, there is no experience of use during pregnancy, other than at birth. Animal experimental studies have shown that epinephrine (adrenaline) has a harmful effect on progeny at doses higher than used for dental anesthesia. After administration by mistake of Orabloc to a blood vessel in the mother, a reduction in blood flow to the uterus may occur as a result of the epinephrine, definedning largering. epinephrine (adrenaline) traction. If you are pregnant, Orabloc 1:200,000 solution for injection should preferably be used instead of Orabloc 1:100,000 solution for

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connection with snort-term treatment was creative.

Driving and using machines

Your dentist will decide when you may drive or use machines again after an intervention. In relevant investigations, no impairment of normal ability to drive was identified after local anaesthetisation with articaine.

Orabloc contains sodium metabisulphite (E223) and sodium.

Sodium metabisulphite (E223) may rarely cause severe hypersensitivity reactions and bronchospasm. This medicinal product

contains less than I mmol sodium (23 mg) per dose, i.e. essentially 'sodium-free'

3. How Orabloc is used

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An injection of 1.8 ml Orabloc per tooth isgenerally enough for the simple removalofuninflamed upper jaw teeth. In a few cases,

a subsequent injection of I - 1.8 ml may be necessary to achieve full local anaesthetisation.

Generally, not every tooth needs to be anaesthetised with the full quantity of Orabloc for the removal of neighbouring teeth. The

number of injections can nonnally be reduced. If an incision or a suture on your palate is necessary, an injection of approx. 0.1 ml per puncture is sufficient. When removing uninflamed front molars in the lower jaw, an injection of 1.8 ml per tooth is generally sufficient. If a full effect does not arise after that, your dentist can give a subsequent injection of 1 - 1.8 ml. Only if the site is not completely anaesthetised even after the subsequent injection your dentist can carry out the otherwise usual anaesthetisation of the entire mandibular nerve. In the case of jaw surgery, your dentist will dose Orabloc I: 100,000 solution for injection individually according to the seriousness and length of the intervention. Adults can receive up to 7 mg articaine per kg body weight in the course of a treatment. Quantities of up to 500 mg(equivalent to 12.5 ml solution for injection) are nonnally tolerated weil. Elderly and patients with severe disturbance offiver and kidney function [n elderly patients and patients with severe disturbance of liver and kidney function(e.g. in the case of nephritis or cirrhosis of the liver), increased quantities of articaine may arise in the blood. If you are in these patient groups, your dentist should take particular care to ensure that the smallest possible quantity for adequated anaesthetisation is used.

care to ensure that the smallest possible quantity for adequate anaesthetisation is used. Use in children and adolescents If Orabloc is used in children and adolescents, the minimum volume necessary to achieve adequate anaesthesia should be used

The injection quantity is to be dosed individually according to the child's and adolescent's age and weight. A maximum dose of 5

The injection quality is a be cused initially accounting to the child said advisement's age and weight. A maximum cuse of any gardicaine per kg body weight should not be exceeded.

This product has not been studied in children less than I year old. Method of administration

Orablo: is intended for administration in the oral cavity (dental use).

To prevent infections (e.g. transmission of hepatitis), new and sterile syringes and needles must be used for each injection. For

single use. Any unused solution should be discarded. If discoloured or cloudy, the medicine must not be used.

- If you received too much of Orabloc sive quantities of Orabloc are used, disturbances of nervous system function may occur, e.g.:

- clouding of consciousness
- seizures and disorders of breathing - seizures and disorders of breatning.

 Disturbances of cardiovascular function may also arise, such as a fall or rise in your blood pressure. Such disturbances require dental monitoring and possibly suitable therapy by your dentist.

 If you have any further questions on the use of this medicine, ask your dentist, doctor or phannacist.

 4. Possible side effects

 Like all medicines, this medicine can cause side effects, although not everybody gets them.

Common (may affect up to 1 in 10 people)

Nausea, vomiting, disturbed sensation of touch (paraesthesia), reduction in sensitivity of sensation in the mouth and facial area

Nausea, vontining, disturbed sensation of totuch (paraesthesia), reduction in sensitivity of sensation in the mouth and facial area (hypoaesthesia); headaches which are presumably attributable to the epinephrine fraction <u>Uncommon</u> (may affect up to 1 in 100 people)
Rapid heart beat, (tachtycardia), dizziness
<u>Uncommon</u> (may affect up to 2 in 100 people)
Rapid heart beat, (tachtycardia), dizziness
<u>Uncommon</u> (may affect up to 2 in 100 people)
Rapid heart facility this disturbances, increase in blood pressure, drop in blood pressure, lowered heart rate (bradycardia), heart failure

- and shock (possibly life-threatening). - Depending ondose (particularly in the case of excessive dosing or inadvertent injection into ablood vessel), disturbances of the
- nervous system may arise, e.g.:

- aguaturi, nervousness, drowsiness ranging to loss of consciousness, coma, respiratory disturbances ranging to respiratory arrest, muscle trenor, muscle witching ranging to convulsions. During or shortly after the injection officeal anaesthetics in the head area, temporary visual disorders (blurred vision, blindness, - Nerve damage(e.g. of the facial nerve) and a reduction insensitivity of taste in the mouth and facial area are not side effects that
- can be induced by Orabloc alone. These side effects mayarise in any dental intervention and cannot therefore be ruledout. They are detennined by the course of the nerves in the injection area or by defective injection technique.

 An inadequate supply ofoxygen to tissue leading to the death of tissue as a result of inadvertent injection into ablood vessel may very rarely arise in the injection area.
- Hypersensitivity reactions (allergic or allergy-like reactions) may arise. These may manifest themselves as swelling or
- inflammation at the injection site. Hypersensitivity reactions arise that are not restricted to the injection site.
- itching,
 connective tissue inflammation,
- Continuo Colo, Facial swelling (in the fon11 ofQuincke's oedema) with swelling of the upper and/or lower lip and/or cheeks, swelling of the laryngeal area with a feeling of tightness and swallowing complaints,
- nettle rash,
- spiratory complaints ranging to anaphylactic shock. Owing to the content of sodium metabisulphite hypersensitivity reactions that may manifest themselves as vomiting, diarrhoea
- ing, acute asthma attacks, disturbances of consciousness or shock may arise very rarely, particularly in bronchia asunitations.

 - respiratory dysfunction (tachypnea, bradypnea) that may lead to apnea

 Additional side effects in children

 In young children, compared to adults, there is an increased risk of self-biting, whichcould result in soft tissue injury, because of

ied numbness in the area of the mouth after the dental procedure.

In a side effect arises suddenly or develops strongly, inform a doctor immediately. This is particularly important because certain pharmaceutical side effects (e.g. fall in blood pressure or respiratory disturbances) may become life-threatening.

Preparting of side effects
If you get any side effects, talk to your doctor, dentist or phannacist. This includes any possible side effects not listed in this leaflet.

By reporting side effects youcan help provide more informationon the safety of this medicine.

5. How to store Orabloc Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton and on the cartridge after «EXP». The expiry date refers to the last day of that month

Teles to the last use of the Internation.

Do not store above 25°C. Store in the original package in order to protect from light.

Do not throw away any medicines via wastewater or household waste. Ask your phannacist how to throw away medicines you no longer use. These measures will help to protect the environment.

6. Contents of the pack and other information

What Orabloc contains

- The other ingredients are:

The active substances are articaine hydrochloride and epinephrine (adrenaline)

Articaine hydrochloride 4% with adrenaline (epinephrine) solution for injection I: 100,000:

1 ml solution for injection contains 40 mg ofarticaine hydrochloride and 0.01 mg epinephrine (adrenaline) as adrenaline tartrate

One cartridge of 1.8 ml of solution for injection contains 72 mg articaine hydrochloride and 0.018 mg epinephrine (adrenaline

isulphite (E223), sodium chloride, hydrochloric acid 2% (for pH adjustment), water for injection. What Orabloc looks like and contents of the pack Trial of control in the ania contents of the pack.

Orabloc is a sheline, clear, colourless solution for injection in clear glass cartridges closed at one end with a bromobutylic rubber plunger and at the other with an aluminium cap and rubber seal.

The cartidge is available in different packages:
The cartridges are packaged in blisters (10 cartridges/blister); the blisters are packaged in a cardboard box containing 5 x 10

Each cartridge is assembled in a plastic injector; each injector containing a cartridge is placed in a sealed blister; the injectors are packaged in a cardboard box together with an Instructions for use of the injector: 50 or 100 units per commercial pack

This medicinal product is authorised in the Member States of the EEA under the following names:

Germany Articainhydrochlorid mit Epinephrin Pierrel 40 mg/ml + 0,01 mg/ml Injektionslösung Austria Orabioc 40 mg/ml + 10 Mikrogramm/ml Injektionslösung France Orabioc 40 mg/ml Adrenalinee au 1/100 000 Solution injectable

Poland Orabloc 40 mg/ml + 0,01 mg/ml United Kingdom Orabloc 1:100,000

Marketing Authorisation Holder and Manufacture

ada Statale Appia, 7 BIS 46/48 – 81043 Capua (CE)