Summary of Product Characteristics

1. NAME OF THE MEDICINAL PRODUCT

BCG-medac, powder and solvent for intravesical suspension

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

After reconstitution, one vial contains: BCG (Bacillus Calmette-Guérin) bacteria derived from Mycobacterium bovis, seed RIVM derived from seed 1173-P22 x 10⁸ to 3 x 10⁹ viable units

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Powder and solvent for intravesical suspension

Powder: White or almost white powder or porous cake with shades of yellow and grey Solvent: Colourless, clear solution

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Treatment of non-invasive urothelial bladder carcinoma:

- curative treatment of carcinoma in situ
- prophylactic treatment of recurrence of:
 - urothelial carcinoma limited to mucosa:
 - Ta G1-G2 if multifocal and/or recurrent tumour
 - Ta G3
 - urothelial carcinoma in lamina propria but not the muscular of the bladder (T1)
 - carcinoma in situ

4.2 Posology and method of administration

BCG-medac must be administered by physicians experienced in this therapy.

BCG-medac is intended for intravesical use following reconstitution. For instructions on reconstitution of the BCG-medac suspension before administration, see section 6.6.

Posology

Adults and the elderly

The content of one vial, reconstituted and diluted as indicated, is required for one instillation into the urinary bladder.

Induction therapy

BCG-therapy should begin about 2-3 weeks after transurethral resection (TUR) or bladder biopsy, and without traumatic catheterisation, and be repeated at weekly intervals for 6 weeks. In intermediate- and high-risk tumours this should be followed by maintenance therapy. Maintenance treatment schemes are described below.

Maintenance therapy

Based on clinical studies, maintenance therapy following induction is highly recommended. The recommended maintenance scheme consists of 3 instillations at weekly intervals given for a minimum of 1 year up to 3 years at month 3, 6, 12, 18, 24, 30, and 36. In this scheme, up to 27 instillations are administered during a period of three years.

Although maintenance therapy reduces recurrence and may reduce progression, the adverse reactions and discomfort of the treatment may outweigh the benefits for some patients. Thus, benefit-risk assessment and consideration of patient preferences is important before beginning or continuing maintenance treatment. The need for maintenance treatment every 6 months beyond the first year of treatment should further be evaluated on the basis of tumour classification and clinical response.

Paediatric population

The safety and efficacy of BCG-medac in children have not been established. No data are available.

Method of administration

The patient should not drink over a period of 4 hours before the instillation until 2 hours after the instillation.

A urethral catheter is inserted into the bladder under aseptic conditions. A sufficient quantity of lubricant should be used to reduce the chance of traumatising the urinary mucosa and therefore the risk of severe complications, and also to reduce discomfort for the patient associated with the procedure. The bladder must be emptied before BCG-instillation. Complete draining of the bladder after catheterisation reduces residual lubricant which may have reached the bladder before BCG-medac is instilled.

BCG-medac is introduced into the bladder by means of a catheter and at low pressure. The instilled BCG-medac suspension must remain in the bladder for a period of 2 hours if possible. During this period the suspension should have sufficient contact with the entire mucosal surface of the bladder. Therefore, the patient should be mobilised as much as possible or, in case of a bed-ridden patient, should be turned over from back to abdomen and vice versa every 15 minutes. After 2 hours, the patient should void the instilled suspension in a sitting position.

In case of no specific medical contraindication, hyperhydratation of the patient is recommended for the 48 hours following each instillation.

Patients treated with BCG-medac should be given the package leaflet and the patient alert card.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

BCG-medac should not be used in immunosuppressed patients or persons with congenital or acquired immune deficiencies, whether due to concurrent disease (e.g. positive HIV serology, leukaemia, lymphoma), cancer therapy (e.g. cytostatic medicinal products, radiation) or immunosuppressive therapy (e.g. corticosteroids).

BCG-medac should not be administered to persons with <u>active</u> tuberculosis. The risk of active tuberculosis must be ruled out by appropriate anamnesis and, if indicated, by diagnostic tests according to local guidelines.

Past history of radiotherapy of the bladder.

Treatment with BCG-medac is contraindicated in women during lactation (see section 4.6).

BCG-medac must not be instilled before 2 to 3 weeks after a TUR, a bladder biopsy or a traumatic catheterisation.

Perforation of the bladder which might result in an increased risk of severe systemic infections (see section 4.4).

<u>Acute</u> urinary tract infection (see section 4.4). Asymptomatic, isolated leukocyturia and asymptomatic bacteriuria are not contraindications for intravesical therapy with BCG-medac, and antibiotic prophylaxis is not necessary.

4.4 Special warnings and precautions for use

BCG-medac must not be used for subcutaneous, intradermal, intramuscular or intravenous administration or vaccination.

Treatment of symptoms, signs or syndrome

See section 4.8.

Handling precautions

BCG-medac should not be handled either in the same room or by the same personnel preparing cytotoxic medicinal products for intravenous administration. BCG-medac should not be handled by a person who presents with well-known immunodeficiency. Contact of BCG-medac with skin and mucosa should be avoided. Contamination can lead to hypersensitivity reaction or infection of the concerned area.

Spillage of BCG-medac

Spillage of BCG-medac suspension should be treated with a disinfectant with proven activity against mycobacteria. Spillage on the skin should be treated with an appropriate disinfectant.

General hygiene for the patient

It is recommended to wash hands and genital area after micturition. This applies especially to the first micturitions following BCG-instillation. If skin lesions are contaminated, the use of an appropriate disinfectant is recommended.

Tuberculin tests

Cutaneous tests

The intravesical treatment with BCG-medac could induce sensitivity to tuberculin and complicate subsequent interpretation of tuberculin cutaneous tests for mycobacterial infection diagnosis. Therefore, reactivity to tuberculin should be measured before administration of BCG-medac.

Detection of Bacillus Calmette-Guérin

Physicians should be aware that a negative germ biopsy and negative test results do not rule out a systemic BCG-infection. In several cases germ detection was not successful even though the patient experienced a systemic BCG-infection. The available methods (microscopy, PCR and/or cultures and/or a detection of tuberculosis-compatible histology) are not reliable.

Severe systemic BCG-infections/reactions

Traumatic instillation could promote BCG-septicaemic events with possible septic shock and a life-threatening situation. For treatment options see section 4.8

Urinary tract infection should be excluded before each bladder instillation of BCG (bladder mucous membrane inflammation may increase the risk of haematological dissemination of BCG). If a urinary tract infection is diagnosed during BCG-therapy, the therapy should be interrupted until the urinalysis is normalised and treatment with antibiotics is completed.

The possibility of severe systemic BCG-infections with the necessity of anti-tuberculosis therapy has to be considered before starting the BCG-therapy, especially in elderly patients (see Elderly patients) and patients with hepatic impairment.

Severe systemic BCG-infections/reactions have been reported in less than 5%. For signs and symptoms please refer to section 4.8.

In case of a suspicion of a systemic infection a physician specialised in infectious diseases should be consulted. BCG-infection can be potentially fatal. For further information please refer to section 4.8.

In contrast to systemic infections, Reiter's syndrome presents as a mainly immuno-mediated reaction, which is not necessarily caused by disseminated BCG but could also be triggered by BCG only localised in the urinary tract system.

Fever or gross haematuria

Treatment should be postponed until resolution of concurrent fever or gross haematuria.

Low bladder capacity

The risk of bladder contracture may increase in patients with low bladder capacity.

HLA-B27

Patients with positive HLA-B27 could have an increase of the occurrence of reactive arthritis or Reiter's syndrome.

Flare-up of latent BCG-infection (including delayed diagnosis)

There have been single case reports in which BCG-bacteria persisted in the body for several years. Those latent BCG-infections might flare-up years after the initial infection, arising especially from granulomatous pneumonitis, abscesses, infected aneurysms, infection of an implant, graft or the surrounding tissue.

The patient has to be made aware of the possibility of late flare-up of latent BCG-infections and educated regarding the actions if symptoms like fever and weight loss of unknown origin occur. In case of suspicion of a flare-up of latent BCG-infection a physician specialised in infectious diseases should be consulted.

Elderly patients

BCG-administration in elderly patients is not contraindicated. However, the risk of a systemic BCG-infection/reaction should be considered before the first administration is performed. Elderly patients may suffer from renal or hepatic impairment which could have an influence on the treatment with anti-tuberculosis medicinal products in case of severe systemic BCG-infection/reaction. Particular caution should also be exercised in elderly patients with reduced general condition.

Pregnancy

BCG-medac is not recommended during pregnancy (see section 4.6).

Patients with contact to immunosuppressed persons

Patients treated with BCG-medac should employ adequate hygienic measures if in contact with immunosuppressed patients. *M. bovis* is less pathogenic than *M. tuberculosis* and man-to-man transmission has not been reported yet, but it cannot be excluded especially in immunosuppressed patients.

Sexual transmission

Sexual transmission of BCG has not been reported yet, but it is recommended to use a condom during coitus for one week after BCG-therapy.

Traceability

In order to improve the traceability of biological medicinal products, the name and the batch number of the administered product should be clearly recorded.

4.5 Interaction with other medicinal products and other forms of interactions

BCG bacteria are sensitive to anti-tuberculous medicinal products (e.g. ethambutol, streptomycin, p-aminosalicylic acid [PAS], isoniazid [INH] and rifampicin), antibiotics and antiseptics. A resistance against pyrazinamide and cycloserine has been described.

During intravesical BCG-instillation therapy, simultaneous administration of anti-tuberculous agents and antibiotics like fluoroquinolones, doxycycline or gentamicin should be avoided due to sensitivity of BCG to those medicinal products.

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no or limited amount of data from the use of BCG in pregnant women. Reproductive animal studies were not performed. BCG-medac is not recommended during pregnancy.

Breast-feeding

There is insufficient information on the excretion of BCG/metabolites in human milk. BCG-medac is contraindicated during breast-feeding (see section 4.3).

Fertility

Intravesical BCG-therapy was found to adversely affect spermatogenesis and might cause oligospermia or azoospermia. Animal studies suggest that these effects might be transient and reversible. However, men should seek advice about the possibility of sperm preservation before starting therapy.

4.7 Effects on ability to drive and use machines

Local or systemic symptoms during therapy with BCG-medac could affect the ability to drive or operate machines.

4.8 Undesirable effects

Undesirable effects are listed below by system organ class and frequency. Frequencies below are defined as: Very common ($\geq 1/10$), common ($\geq 1/100$ to < 1/10), uncommon ($\geq 1/1,000$ to < 1/100), rare ($\geq 1/10,000$ to < 1/1,000), very rare (< 1/10,000) or not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

System organ class	Frequency and undesirable effects
Infections and	Very common
infestations	Cystitis and inflammatory reactions (granuloma) of the bladder,
	asymptomatic granulomatous prostatitis
	Uncommon
	Urinary tract infection, orchitis, epididymitis, symptomatic
	granulomatous prostatitis, severe systemic
	BCG-reaction/infection, BCG-sepsis, miliary pneumonitis, skin
	abscess, Reiter's syndrome (conjunctivitis, asymmetrical
	oligoarthritis and cystitis)
	Rare
	Vascular infection (e.g. infected aneurysm), renal abscess
	Very rare
	BCG-infection of implants and surrounding tissue (e.g. aortic
	graft infection, cardiac defibrillator, hip or knee arthroplasty),
	regional lymph node infection, osteomyelitis, bone marrow
	infection, peritonitis, psoas abscess, infection of the glans penis,
	orchitis or epididymitis resistant to anti-tuberculous therapy
Blood and lymphatic	Uncommon
system disorders	Cytopenia, anaemia
5 y 5 w 111 u 1501 u 15	Very rare
	Cervical lymphadenitis
	Not known
	Haemophagocytic syndrome
Immuno avatom	
Immune system disorders	Very common Transient systemic PCC respire (favor < 28.5 °C, fly like
uisoiueis	Transient systemic BCG-reaction (fever < 38.5 °C, flu-like
	symptoms including malaise, fever, chills, general discomfort, myalgia)
	<u>Very rare</u> <u>Hypersonaitivity reaction (a.g. codema of evolide, cough)</u>
Eva disordara	Hypersensitivity reaction (e.g. oedema of eyelids, cough)
Eye disorders	<u>Very rare</u> Chorioretinitis, conjunctivitis, uveitis
Vascular disorders	
v asculat utsoluers	Uncommon
	Hypotension Very rare
	Very rare Vegender fistule
	Vascular fistula
	Not known Vacculitie (including vacculitie of the central nervous system)
Descinatory (1	Vasculitis (including vasculitis of the central nervous system)
Respiratory, thoracic	<u>Uncommon</u>
and mediastinal	Pulmonary granuloma
disorders	
Gastrointestinal	Very common
disorders	Nausea
	Common
	Diarrhoea, abdominal pain
	Very rare
	Vomiting, intestinal fistula
Hepatobiliary disorders	<u>Uncommon</u>
	Hepatitis
Skin and subcutaneous	Uncommon
Similand Subcutaneous	
	Skin rash
tissue disorders Musculoskeletal and	Skin rash Common

disorders	Uncommon	
	Arthritis, arthralgia	
Renal and urinary	Very common	
disorders	Frequent urination with discomfort and pain	
	Common	
	Urinary incontinence	
	Uncommon	
	Macroscopic haematuria, bladder retention, urinary tract	
	obstruction, contracted bladder	
	Not known	
	Renal failure, pyelonephritis, nephritis (including	
	tubulointerstitial nephritis, interstitial nephritis and	
	glomerulonephritis)	
Reproductive system	Not known	
and breast disorders	Genital disorders (e.g. vaginal pain, dyspareunia), oligospermia,	
	azoospermia	
General disorders and	<u>Very common</u>	
administration site	Fatigue	
conditions	Common	
	Fever $> 38.5 $ °C	
	<u>Very rare</u>	
	Peripheral Oedema	
Investigations	<u>Uncommon</u>	
	Hepatic enzyme increased	
	<u>Not known</u>	
	Prostatic specific antigen (PSA) increased	

Adverse reactions of BCG-treatment are frequent but generally mild and transient. Adverse reactions usually increase with the number of BCG-instillations.

In common cases myalgia and in uncommon cases, arthritis/arthralgias and skin rash may occur. In most cases of arthritis, arthralgias and skin rash, these can be attributed to hypersensitivity reactions of the patient to BCG. It may be necessary in some cases to discontinue the administration of BCG-medac.

Local adverse reactions

Discomfort and pain when urinating and frequent urination occur in up to 90 % of patients. The cystitis and inflammatory reaction (granulomata) may be an essential part of the anti-tumour activity. Further local adverse reactions which are uncommonly observed: macroscopic haematuria, urinary tract infection, bladder retraction, urinary obstruction, bladder contracture, symptomatic granulomatous prostatitis, orchitis and epididymitis. Renal abscess is rarely observed. Furthermore, genital disorders (e.g. vaginal pain, dyspareunia) may occur with an unknown frequency.

Transient systemic BCG-reaction

Low-grade fever, flu-like symptoms and general discomfort may occur. These symptoms usually subside within 24 - 48 hours and should be managed by standard symptomatic treatment. These reactions are signs of a starting immune reaction. All patients receiving the medicinal product should be carefully monitored and advised to report all incidences of fever and other events outside the urinary tract.

Severe systemic adverse reactions/infections

Distinguishing a BCG-infection from a BCG immune reaction poses a challenge, as the symptoms are very similar at the beginning. In contrast to this a transient systemic BCG-reaction is a very common adverse reaction that must be differentiated.

The clinical signs and symptoms of a BCG-infection/reaction at the beginning are fever > 39.5 °C during at least 12 hours, fever > 38.5 °C during at least 48 hours and worsening of general condition. Typical signs of an infection are the development of miliary pneumonia, granulomatous hepatitis, liver function test abnormalities (especially an elevation in alkaline phosphatase), organic dysfunction (other than genito-urinary tract) with granulomatous inflammation at biopsy over time. In case of a suspicion of a systemic infection a physician specialised in infectious diseases should be consulted. BCG-infection can be potentially fatal.

Although the symptoms of a systemic BCG-infection do not differ from tuberculosis, the patient does not need to be isolated, because *M. bovis* is less pathogenic for humans than *M. tuberculosis*.

<u>In case of a flare-up of a latent infection</u> the patients usually present with symptoms of fever and weight loss of unknown origin. Several case reports show that the diagnosis is challenging as the symptoms vary and a causal relationship with BCG-infection is not suspected by physicians. A correct and early diagnosis and as a consequence, an appropriate treatment is important for the outcome, especially in elderly or debilitated patients, to avoid fatal consequences. Please note that a patient alert card is available with focus on the topic which has to be handed over to the patient (see also section 4.4).

In case of suspicion of a flare-up of latent BCG-infection a physician specialised in infectious diseases should be consulted.

The additional use of corticosteroids might be recommended in case of sepsis, granulomatous reactions (e.g. lung or liver) and other immune-mediated reactions.

Treatment of symptoms, signs and syndrome			
Symptoms, signs or syndrome	Treatment		
1) Symptoms of vesical	Symptomatic treatment		
irritation lasting less than			
48 hours			
2) Symptoms of vesical	Discontinue therapy with BCG-medac and start treatment		
irritation lasting more or equal	with quinolones. If after 10 days no complete resolution is		
to 48 hours	observed, administer isoniazid (INH)* for 3 months.		
	In case of anti-tuberculosis treatment, therapy with		
	BCG-medac should definitively be discontinued.		
3) Concomitant bacterial	Postpone BCG-medac therapy until the urinalysis is		
infection of urinary tract	normalised and treatment with antibiotics is completed.		
4) Other genitourinary	Discontinue therapy with BCG-medac.		
undesirable effects:	Administer isoniazid (INH)* and rifampicin*, for 3 to		
symptomatic granulomatous	6 months according to severity.		
prostatitis, epididymitis and			
orchitis, urethral obstruction	In case of anti-tuberculosis treatment, therapy with		
and renal abscess	BCG-medac should definitively be discontinued.		
5) Fever less than 38.5 °C	Symptomatic treatment with paracetamol.		
lasting less than 48 hours			
6) Cutaneous eruption,	Discontinue therapy with BCG-medac.		
arthralgias or arthritis or	Consider a consultation with a specialist for infectious		
Reiter's syndrome	diseases.		
	Administer antihistaminic or non-steroidal anti-		
	inflammatory drugs. Cortisone therapy should be		

Treatment recommendations see table below:

	considered in case of an immune-mediated reaction. If no response, administer isoniazid* for 3 months.
	In case of anti-tuberculosis treatment, therapy with BCG-medac should definitively be discontinued.
7) Systemic	Definitely discontinue therapy with BCG-medac.
BCG-reaction/infection**	Consider a consultation with a specialist for infectious
without septic shock signs	diseases.
	Administer a triple-drug anti-tuberculosis therapy* for
	6 months and low dose corticosteroid therapy.
8) Systemic	Definitely discontinue treatment with BCG-medac.
BCG-reaction/infection with	Administer immediately a triple anti-tuberculosis therapy*
septic shock signs	combined with high-dose, quick-acting corticosteroids.
	Seek the opinion of a specialist for infectious diseases.

*Caution: BCG-bacteria are sensitive to all anti-tuberculous medicinal products currently used, except for pyrazinamide. If a triple anti-tuberculosis therapy is necessary, the combination usually recommended is isoniazid (INH), rifampicin and ethambutol. ** definition see above

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via HPRA Pharmacovigilance, Website: <u>www.hpra.ie.</u>

4.9 Overdose

Overdose is unlikely to occur as one vial of BCG-medac corresponds to one dose.

There are no data indicating that an overdose may lead to any other symptoms than the described undesirable effects.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Immunostimulating agent, ATC code: L03AX03

BCG-medac is a lyophilised suspension of live Bacillus Calmette-Guérin bacteria with low infectious potential derived from *Mycobacterium bovis*, strain RIVM.

Mechanism of action

BCG-medac stimulates the immune system and has anti-tumour activity. Study data suggest that BCG acts as a non-specific immunopotentiator, not by a single mechanism but by a variety of actions involving cells of the immune system. BCG has a stimulating effect on the spleen, enhances macrophage function in the spleen and activates natural killer cells. BCG-instillation stimulates the increase of granulocytes, monocytes/macrophages and T-lymphocytes, indicating local activation of the immune system. Cytokines IL1, IL2, IL6 and TNF α are also increased.

5.2 Pharmacokinetic properties

Most of the bacilli are excreted in the urine in the first hours after the instillation. Whether mycobacteria might be able to pass the intact urothelial wall is still unknown. There have been single case reports in which BCG-bacteria persisted in the urinary tract for more than 16 months (see section 4.4).

5.3 Preclinical safety data

BCG-strain RIVM was tested for toxicity, immunostimulatory properties and anti-tumour activity in a variety of animals. High doses of BCG caused weight retardation in mice, and liver disturbance was also observed. Intravenous injection in rabbits appeared to be pyrogenic. Repeated instillations in guinea pigs induced inflammatory reactions in the bladder wall. As unwanted adverse reactions granulomatous lesions in the liver and lung were observed after high doses. Intravesical application in dogs showed minimal mechanical lesions of the urothelium whereas no signs of active inflammation were observed in the suburothelial stroma.

No mutagenicity, carcinogenicity and reproduction studies have been performed.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Powder: polygeline, glucose anhydrous and polysorbate 80. Solvent: sodium chloride and water for injections.

6.2 Incompatibilities

BCG-medac is incompatible with hypotonic and hypertonic solutions.

6.3 Shelf life

2 years, or 3 years when the amount of viable units at release is greater than 5 x 10^8 cfu/vial, in any case not longer than 4 years from the date of harvest.

After reconstitution the product should be used immediately.

6.4 Special precautions for storage

Store in a refrigerator (2 °C – 8 °C). Do not freeze. Store in the original package in order to protect from light.

For storage conditions after reconstitution of the medicinal product, see section 6.3.

6.5 Nature and contents of container

Powder in a vial (type I glass) with a rubber stopper + 50 ml of solvent in a bag (PVC) with a vial connector and a catheter adapter (conical or Luer-Lock adapter).

Pack sizes (conical adapter): 1, 3, 5 or 6 with or without catheter(s), connector(s). Pack sizes (Luer Lock adapter): 1 or 3 with or without catheter(s).

Not all pack sizes may be marketed.

6.6 Special precautions for disposal and other handling

Instructions for use/handling

Administration of the catheter should be done carefully to avoid injuries of the epithelium which may lead to development of systemic BCG-infection. Use of a lubricant is recommended to minimise the risk of traumatic catheterisation and to reduce patient discomfort. Women might need less lubricant than men. A draining of the bladder after catheterisation reduces residual lubricant before BCG is applied.

Before use, the product has to be resuspended under aseptic conditions with sterile 9 mg/ml (0.9 %) sodium chloride solution (see below). Remix the suspension before use by rotating gently. Avoid skin contact with BCG-medac. The use of gloves is recommended. Visible macroscopic particles do not affect the efficacy and safety of the medicinal product.

For further information regarding the catheter please see the corresponding instructions for use.

The following handling instructions apply to the system with conical or Luer-Lock adapter.



1. Tear open the protective bag but do not remove it completely! This will protect the tip of the instillation system from contamination up to the last minute.



2. Remove the caps from the vial and instillation system. Lay out a disposal bag.



3. Place the BCG-medac vial on a solid surface (e.g. table) and press the vial connector of the instillation system upright and firmly onto the BCG-medac vial. Turn the vial 2 times completely in the same direction.



4. Break open the mechanism in the tube of the vial connector by repeated bending backwards and forwards. This establishes the connection. Please hold the tube – and not the vial – during this process!



5. Pump the liquid into the vial, but do not fill the vial completely. If flow is not possible, turn the vial again 2 times completely in the same direction to assure that the septum is completely penetrated. Repeat this step until flow is possible.



6. Invert the entire system. Pump air from the instillation system into the vial at the top and draw the reconstituted BCG-medac into the instillation system. Do not remove the vial.



7. Keep the instillation system in an upright position. Now remove the protective bag completely. Connect the catheter (and a connector [conical to Luer-Lock]) to the instillation system. Now break the sealing mechanism in the tube section by bending it back and forth and instil the BCGmedac suspension into the patient's bladder. At the end of the instillation free the catheter by pressing air through. Keep the instillation system squeezed and place it together with the catheter into the disposal bag.



Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORISATION HOLDER

medac Gesellschaft für klinische Spezialpräparate mbH Theaterstrasse 6 22880 Wedel Germany

8. MARKETING AUTHORISATION NUMBER(S)

PA0623/004/001

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 05 April 2002 Date of last renewal: 02 October 2006

10. DATE OF REVISION OF THE TEXT

December 2023