SUMMARY OF PRODUCT CHARACTERISTICS

1 NAME OF THE MEDICINAL PRODUCT
Dexamethasone 3.3 mg/ml Solution for Injection or Infusion

2 QUALITATIVE AND QUANTITATIVE COMPOSITION
Each ml of solution contains 3.3 mg dexamethasone (as sodium phosphate) which is equivalent to 4 mg dexamethasone phosphate or 4.37 mg dexamethasone sodium phosphate.
Each 2 ml contains 6.6 mg dexamethasone (as sodium phosphate) which is equivalent to 8 mg dexamethasone phosphate or 8.74 mg dexamethasone sodium phosphate.
For the full list of excipients, see 6.1.

3 PHARMACEUTICAL FORM
Solution for injection or infusion.
Clear, colourless to slightly yellowish solution, having pH ranging from 7.0 to 8.5.

4 CLINICAL PARTICULARS

4.1 Therapeutic indications
Dexamethasone can be used for all forms of general and local glucocorticoid injection therapy and all acute conditions in which intravenous glucocorticoids may be life-saving.

4.2 Posology and method of administration
Dosage

N.B. For this section of document all doses are expressed as mg dexamethasone
In general, glucocorticoid dosage depends on the severity of the condition and response of the patient. Under certain circumstances, for instance in stress, extra dosage adjustments may be necessary. If no favourable response is noted within a couple of days, glucocorticoid therapy should be discontinued.

**Adults and Elderly**

Once the disease is under control the dosage should be reduced or tapered off to the lowest suitable level under continuous monitoring and observation of the patient (See Section 4.4).

For acute life-threatening situations (e.g. anaphylaxis, acute severe asthma) substantially higher dosages may be needed. Cerebral oedema (adults): initial dose 8-16 mg iv followed by 5 mg iv or im every 6 hours, until a satisfactory result has been obtained. In brain surgery these dosages may be necessary until several days after the operation. Thereafter, the dosage has to be tapered off gradually. Increase of intracranial pressure associated with brain tumours can be counteracted by continuous treatment.

For local treatment, the following dosages can be recommended:

- intra-articulary: 1.6-3 mg large joints
  0.6-0.8 mg small joints
- intrabursally: 1.6-3 mg;
- in tendon sheaths: 0.3-0.8mg

The frequency of these injections may vary from every 3-5 days to every 2-3 weeks.

For rectal drip in cases of ulcerative colitis: 4 mg diluted in 120 ml saline.

**Suggested doses for children**

Dosage requirements are variable and may have to be changed according to individual needs. Usually 0.2 mg/kg to 0.4 mg/kg of body weight daily.

**Administration**

Dexamethasone injections may be administered intravenously, subcutaneously, intramuscularly, by local injection or as a rectal drip. For administration by intravenous infusion: see section on compatibility with infusion fluids. With intravenous administration high plasma levels can be obtained rapidly.

Rapid intravenous injection of massive doses of glucocorticoids may sometimes cause cardiovascular collapse; the injection should therefore be given slowly over a period of several minutes.
Intra-articular injections should be given under strictly aseptic conditions.

4.3 Contraindications

Systemic infection unless specific anti-infective therapy is employed.

Hypersensitivity to any ingredient.

Local injection of a glucocorticoid is contraindicated in bacteraemia and systemic fungal infections, unstable joints, infection at the injection site e.g. septic arthritis resulting from gonorrhea or tuberculosis.

4.4 Special warnings and precautions for use

A patient information leaflet should be supplied with this product.

In post-marketing experience tumour lysis syndrome (TLS) has been reported in patients with haematological malignancies following the use of dexamethasone alone or in combination with other chemotherapeutic agents. Patients at high risk of TLS, such as patients with high proliferative rate, high tumour burden, and high sensitivity to cytotoxic agents, should be monitored closely and appropriate precautions taken.

Patients and/or carers should be warned that potentially severe psychiatric adverse reactions may occur with systemic steroids (see section 4.8). Symptoms typically emerge within a few days or weeks of starting the treatment. Risks may be higher with high doses/systemic exposure (see also section 4.5 for pharmacokinetic interactions that can increase the risk of side effects), although dose levels do not allow prediction of the onset, type severity or duration of reactions. Most reactions recover after either dose reduction or withdrawal, although specific treatment may be necessary. Patients/carers should be encouraged to seek medical advice if worrying psychological symptoms develop, especially if depressed mood or suicidal ideation is suspected. Patients/carers should also be alert to possible psychiatric disturbances that may occur either during or immediately after dose tapering/withdrawal of systemic steroids, although such reactions have been reported infrequently.

Particular care is required when considering the use of systemic corticosteroids in patients with existing or previous history of severe affective disorders in themselves or in their first degree relatives. These would include depressive or manic-depressive illness and previous steroid psychosis.

Undesirable effects may be minimised by using the lowest effective dose for the minimum period, and by administering the daily requirement as a single
morning dose or whenever possible as a single morning dose on alternative
days. Frequent patient review is required to appropriately titrate the dose
against disease activity.

After parenteral administration of glucocorticoids serious anaphylactoid
reactions, such as glottis oedema, urticaria and bronchospasm, have
occasionally occurred, particularly in patients with a history of allergy. If such
an anaphylactoid reaction occurs, the following measures are recommended:
 immediate slow intravenous injection of 0.1 - 0.5 ml of adrenaline (solution of
1:1000: 0.1 - 0.5 mg adrenaline dependent on body weight), intravenous
administration of aminophylline and artificial respiration if necessary.

Corticosteroids should not be used for the management of head injury or
stroke because it is unlikely to be of any benefit and may even be harmful.

The results of a randomised, placebo-controlled study suggest an increase in
mortality if methylprednisolone therapy starts more than two weeks after the
onset of Acute Respiratory Distress Syndrome (ARDS). Therefore, treatment
of ARDS with corticosteroids should be initiated within the first two weeks of
onset of ARDS (See also section 4.2).

**Preterm neonates:**

Available evidence suggests long-term neurodevelopmental adverse events
after early treatment (< 96 hours) of premature infants with chronic lung
disease at starting doses of 0.25 mg/kg twice daily.

**Dexamethasone withdrawal**

Adrenal cortical atrophy develops during prolonged therapy and may persist
for years after stopping treatment. Withdrawal of corticosteroids after
prolonged therapy must therefore always be gradual to avoid acute adrenal
insufficiency, being tapered off over weeks or months according to the dose
and duration of treatment.

In patients who have received more than physiological doses of systemic
corticosteroids (approximately 1 mg dexamethasone) for greater than 3 weeks,
withdrawal should not be abrupt. How dose reduction should be carried out
depends largely on whether the disease is likely to relapse as the dose of
systemic corticosteroids is reduced. Clinical assessment of disease activity
may be needed during withdrawal. If the disease is unlikely to relapse on
withdrawal of systemic corticosteroids but there is uncertainty about HPA
suppression, the dose of systemic corticosteroid may be reduced rapidly to
physiological doses. Once a daily dose of 1mg dexamethasone is reached, dose
reduction should be slower to allow the HPA-axis to recover.

Abrupt withdrawal of systemic corticosteroid treatment, which has continued
up to 3 weeks is appropriate if it is considered that the disease is unlikely to
relapse. Abrupt withdrawal of doses of up to 6mg daily of dexamethasone for
3 weeks is unlikely to lead to clinically relevant HPA-axis suppression in the
majority of patients. In the following patient groups, gradual withdrawal of systemic corticosteroid therapy should be considered even after courses lasting 3 weeks or less:

- Patients who have had repeated courses of systemic corticosteroids, particularly if taken for greater than 3 weeks.
- When a short course has been prescribed within one year of cessation of long-term therapy (months or years).
- Patients who may have reasons for adrenocortical insufficiency other than exogenous corticosteroid therapy.
- Patients receiving doses of systemic corticosteroid greater than 6mg daily of dexamethasone.
- Patients repeatedly taking doses in the evening.

During prolonged therapy any inter-current illness, trauma or surgical procedure will require a temporary increase in dosage; if corticosteroids have been stopped following prolonged therapy they may need to be temporarily re-introduced.

Patients should carry 'Steroid treatment' cards which give clear guidance on the precautions to be taken to minimise risk and which provide details of prescriber, drug, dosage and the duration of treatment.

Anti-inflammatory/Immunosuppressive effects and Infection

Suppression of the inflammatory response and immune function increases the susceptibility to infections and their severity. The clinical presentation may often be atypical, and serious infections such as septicaemia and tuberculosis may be masked and may reach an advanced stage before being recognised.

Appropriate antimicrobial therapy should accompany glucocorticoid therapy when necessary e.g. in tuberculosis and viral and fungal infections of the eye.

*Chickenpox is of particular concern since this normally minor illness may be fatal in immunosuppressed patients.* Patients (or parents of children) without a definite history of chickenpox should be advised to avoid close personal contact with chickenpox or herpes zoster and if exposed they should seek urgent medical attention. Passive immunisation with varicella zoster immunoglobulin (VZIG) is needed by exposed non-immune patients who are receiving systemic corticosteroids or who have used them within the previous 3 months; this should be given within 10 days of exposure to chickenpox. If a diagnosis of chickenpox is confirmed, the illness warrants specialist care and urgent treatment. Corticosteroids should not be stopped and the dose may need to be increased.
Measles. Patients should be advised to take particular care to avoid exposure to measles and to seek immediate medical advice if exposure occurs; prophylaxis with intramuscular normal immunoglobin may be needed.

Live vaccines should not be given to individuals with impaired immune responsiveness. The antibody response to other vaccines may be diminished.

Special precautions

Particular care is required when considering the use of systemic corticosteroids in patients with the following conditions and frequent patient monitoring is necessary.

a. Osteoporosis (post-menopausal females are particularly at risk).

b. Hypertension or congestive heart failure.

c. Existing or previous history of severe affective disorders (especially previous steroid psychosis).

d. Diabetes mellitus (or a family history of diabetes).

e. History of tuberculosis, since glucocorticoids may induce reactivation.

f. Glaucoma (or a family history of glaucoma).

g. Previous corticosteroid-induced myopathy.

h. Liver failure.

i. Renal insufficiency.

j. Epilepsy.

k. Gastro-intestinal ulceration.

l. Migraine

m. Certain parasitic infestations in particular amoebiasis.

n. Incomplete statural growth since glucocorticoids on prolonged administration may accelerate epiphyseal closure.

o. Patients with Cushing's syndrome

In the treatment of conditions such as tendinitis or tenosynovitis care should be taken to inject into the space between the tendon sheath and the tendon as cases of ruptured tendon have been reported.

Use in children
Corticosteroids cause dose-related growth retardation in infancy, childhood and adolescence, which may be irreversible.

Dexamethasone has been used 'off label' to treat and prevent chronic lung disease in preterm infants. Clinical trials have shown a short term benefit in reducing ventilator dependence but no long term benefit in reducing time to discharge, the incidence of chronic lung disease or mortality. Recent trials have suggested an association between the use of dexamethasone in preterm infants and the development of cerebral palsy. In view of this possible safety concern, an assessment of the risk:benefit should be made on an individual patient basis.

**Use in the Elderly**

The common adverse effects of systemic corticosteroids may be associated with more serious consequences in old age, especially osteoporosis, hypertension, hypokalaemia, diabetes, susceptibility to infection and thinning of the skin. Close clinical supervision is required to avoid life-threatening reactions.

**Excipients**

This medicine contains 0.4 mg of sodium per 1 ml ampoule and 0.8 mg of sodium per 2 ml ampoule (less than 23 mg per ampoule), i.e. it is essentially sodium free.

**4.5 Interaction with other medicinal products and other forms of interaction**

Rifampicin, rifabutin, ephedrine, carbamazepine, phenylbutazone, phenobarbital, phenytoin, primidone, and aminoglutethimide enhance the metabolism of corticosteroids and its therapeutic effects may be reduced.

The effects of anticholinesterases are antagonised by corticosteroids in myasthenia gravis.

The desired effects of hypoglycaemic agents (including insulin), anti-hypertensives, cardiac glycosides and diuretics are antagonised by corticosteroids, and the hypokalaemic effects of acetazolamide, loop diuretics, thiazide diuretics and carbenoxolone are enhanced.

The efficacy of coumarin anticoagulants may be enhanced by concurrent corticosteroid therapy and close monitoring of the INR or prothrombin time is required to avoid spontaneous bleeding.

The renal clearance of salicylates is increased by corticosteroids and steroid withdrawal may result in salicylate intoxication. There may be interaction with salicylates in patients with hypoprothrombinaemia.

Co-treatment with CYP3A inhibitors, including cobicistat-containing products, is expected to increase the risk of systemic side-effects. The combination
should be avoided unless the benefit outweighs the increased risk of systemic corticosteroid side-effects, in which case patients should be monitored for systemic corticosteroid side effects.

4.6  **Fertility, pregnancy and lactation**

**Pregnancy**

The ability of corticosteroids to cross the placenta varies between individual drugs, however, dexamethasone readily crosses the placenta.

Administration of corticosteroids to pregnant animals can cause abnormalities of foetal development including cleft palate, intra-uterine growth retardation and affects on brain growth and development. There is no evidence that corticosteroids result in an increased incidence of congenital abnormalities, such as cleft palate/lip in man (see also section 5.3). However, when administered for prolonged periods or repeatedly during pregnancy, corticosteroids may increase the risk of intra-uterine growth retardation. Hypoadrenalism may, in theory, occur in the neonate following prenatal exposure to corticosteroids but usually resolves spontaneously following birth and is rarely clinically important. As with all drugs, corticosteroids should only be prescribed when the benefits to the mother and child outweigh the risks. When corticosteroids are essential however, patients with normal pregnancies may be treated as though they were in the non-gravid state.

**Lactation**

Corticosteroids may pass into breast milk, although no data are available for dexamethasone. Infants of mothers taking high doses of systemic corticosteroids for prolonged periods may have a degree of adrenal suppression.

4.7  **Effects on ability to drive and use machines**

None known

4.8  **Undesirable effects**

**Side-effects**

Local adverse reactions include post-injection flare, and a painless destruction of the joint reminiscent of Charcots arthropathy especially with repeated intra-articular injection.

The incidence of predictable undesirable effects, including hypothalamic-pituitary-adrenal suppression correlates with the relative potency of the drug,
dosage, timing of administration and the duration of treatment. Cases of ruptured tendon have been reported (see Section 4.4).

Local injection of glucocorticoid may produce systemic effects.

**Endocrine/metabolic**


**Anti-inflammatory and Immunosuppressive effects**

Increased susceptibility and severity of infections with suppression of clinical symptoms and signs. Diminished lymphoid tissue and immune response. Opportunistic infections, recurrence of dormant tuberculosis and decreased responsiveness to vaccination and skin tests. (see Section 4.4).

**Musculoskeletal**

Osteoporosis, vertebral and long bone fractures, avascular osteonecrosis, tendon rupture.

Proximal myopathy.

**Fluid and electrolyte disturbance**

Sodium and water retention, hypertension, potassium loss, hypokalaemic alkalosis.

**Neuropsychiatric**

A wide range of psychiatric reactions including affective disorders (such as irritable, euphoric, depressed and labile mood, and suicidal thoughts), psychotic reactions (including mania, delusions, hallucinations, and aggravation of schizophrenia), behavioural disturbances, irritability, anxiety, sleep disturbances, and cognitive dysfunction including confusion and amnesia have been reported. Reactions are common and may occur in both adults and children. In adults, the frequency of severe reactions has been estimated to be 5-6%. Psychological effects have been reported on withdrawal of corticosteroids; the frequency is unknown.


**Ophthalmic**
Increased intra-ocular pressure, glaucoma, papilloedema, posterior subcapsular cataracts, corneal or scleral thinning, exacerbation of ophthalmic viral or fungal diseases.

**Not known:** frequency cannot be estimated from the available data

Chorioretinopathy

**Gastrointestinal**

Dyspepsia, peptic ulceration with perforation and haemorrhage, acute pancreatitis, candidiasis.

**Dermatological**

Impaired healing, skin atrophy, bruising, telangiectasia, striae, increased sweating and acne.

**General**

Hypersensitivity including anaphylaxis, has been reported. Leucocytosis. Thromboembolism.

A transient burning or tingling sensation mainly in the perineal area following intravenous injection of large doses of corticosteroid phosphates.

**Withdrawal symptoms and signs**

Too rapid a reduction of corticosteroid dosage following prolonged treatment can lead to acute adrenal insufficiency, hypotension and death. (see Section 4.4).

A 'withdrawal syndrome' may also occur including, fever, myalgia, arthralgia, rhinitis, conjunctivitis, painful itchy skin nodules and loss of weight.

**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 the Yellow Card Scheme, Website: www.mhra.gov.uk/yellowcard.

### 4.9 Overdose

It is difficult to define an excessive dose of a corticosteroid as the therapeutic dose will vary according to the indication and patient requirements. Massive iv corticosteroid doses given as a pulse in emergencies are relatively free from hazardous effects.
Exaggeration of corticosteroid related adverse effects may occur. Treatment should be asymptomatic and supportive as necessary.

5 PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties
Dexamethasone is a synthetic adrenocorticoid with approximately a 7 times higher anti-inflammatory potency than prednisolone and 30 times that of hydrocortisone. Adrenocorticoids act on the HPA at specific receptors on the plasma membrane. On other tissues the adrenocorticoids diffuse across cell membranes and complex with specific cytoplasmic receptors which enter the cell nucleus and stimulate protein synthesis. Adrenocorticoids have anti-allergic, antitoxic, antishock, antipyretic and immunosuppressive properties. Dexamethasone has only minor mineralocorticoid activities and does therefore, not induce water and sodium retention.

5.2 Pharmacokinetic properties
After administration of Dexamethasone Solution for Injection or Infusion, dexamethasone sodium phosphate is rapidly hydrolysed to dexamethasone. After an iv dose of 20mg dexamethasone, plasma levels peak within 5 minutes. Dexamethasone is bound (up to 77%) by plasma proteins, mainly albumin. There is a high uptake of dexamethasone by the liver, kidney and adrenal glands. Metabolism in the liver is slow and excretion is mainly in the urine, largely as unconjugated steroids. The plasma half life is 3.5-4.5 hours but as the effects outlast the significant plasma concentrations of steroids the plasma half-life is of little relevance and the use of biological half life is more applicable. The biological half life of dexamethasone is 36-54 hours, therefore dexamethasone is especially suitable in conditions where continuous glucocorticoid action is desirable.

5.3 Preclinical safety data
In animal studies, cleft palate was observed in rats, mice, hamsters, rabbits, dogs and primates; not in horses and sheep. In some cases these divergences were combined with defects of the central nervous system and of the heart. In primates effects in the brain were seen after exposure. Moreover, intra-uterine growth can be delayed. All these effects were seen at high dosages.
6 PHARMACEUTICAL PARTICULARS

6.1 List of excipients
Creatinine
Absorbic Acid (E300)
Sodium Citrate (E331)
Water for injections
Sodium hydroxide (E524) (for pH adjustment)

6.2 Incompatibilities
Dexamethasone sodium phosphate is physically incompatible with
daunorubicin, doxorubicin and vancomycin and should not be admixed with
solutions containing these drugs. Also incompatible with doxapram HCl and
glycopyrrolate in syringe.

6.3 Shelf life
21 months

Chemical and physical in-use stability has been demonstrated for 24 h at
room temperature and in daylight conditions when diluted with the infusion
fluids listed in 6.6

From a microbiological point of view, the product should be used
immediately. If not used immediately, in-use storage timer and conditions
prior to use are the responsibility of the user and would normally not be longer
than 24 hours at 2-8°C, unless dilution has taken place in controlled and
validated aseptic conditions.

6.4 Special precautions for storage
Do not store above 25°C. Do not refrigerate or freeze.
Store in the original package in order to protect from light.
For storage conditions after dilution of the medicinal product, see section 6.3.

6.5 Nature and contents of container
Colourless, neutral, type I glass ampoules.
1ml glass ampoules in packs of 5 or 10.
2ml glass ampoules in packs of 5.
Not all pack sizes may be marketed.

6.6 **Special precautions for disposal**

*Use with infusion fluids*

Dexamethasone can be diluted with the following infusion fluids:

- sodium chloride 0.9%
- anhydrous glucose 5%
- invert sugar 10%
- sorbitol 5%
- ringer's solution
- ringer-lactate
- dextran 40 10% w/v

Using these infusion fluids, Dexamethasone Injection can also be injected into the infusion line without causing precipitation of the ingredients. Direct injection into the infusion line is also possible with mannitol 10%.

For single use only.

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

7 **MARKETING AUTHORISATION HOLDER**

Wockhardt UK Ltd, Ash Road North, Wrexham LL13 9UF, UK

8 **MARKETING AUTHORISATION NUMBER(S)**

PL 29831/0667

9 **DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION**

15/01/2015

10 **DATE OF REVISION OF THE TEXT**

07/03/2017