1	AUGMENTIN BD S
2	AUGMENTIN BD SF
3	Amoxicillin trihydrate – Potassium clavulanate
4	
5	QUALITATIVE AND QUANTITATIVE COMPOSITION
6 7 8	AUGMENTIN BD S suspension L: When reconstituted each 5 mL contains 200 mg amoxicillin (as amoxicillin trihydrate) and 28.5 mg clavulanic acid (as potassium clavulanate).
9 10 11	AUGMENTIN BD SF suspension: When reconstituted each 5 mL contains 400 mg amoxicillin (as amoxicillin trihydrate) and 57 mg clavulanic acid (as potassium clavulanate).
12 13	Sugar-free. Contains sweetener (aspartame 12.5 mg/5 ml).
14	PHARMACEUTICAL FORM
15 16	A white to off-white dry powder for reconstitution in water to form an off-white mixed-fruit flavoured suspension.
17	CLINICAL PARTICULARS
18	Indications
19 20	AUGMENTIN should be used in accordance with local official antibiotic-prescribing guidelines and local susceptibility data.
21 22 23 24	AUGMENTIN BD S and AUGMENTIN BD SF, for twice daily oral dosing, is indicated for short term treatment of bacterial infections at the following sites when amoxicillin resistant beta-lactamase producing strains are suspected as the cause. In other situations, amoxicillin alone should be considered.
25 26	Upper respiratory tract infections (including ENT) e.g. recurrent tonsillitis, sinusitis, otitis media.
27 28	Lower respiratory tract infections e.g. acute exacerbations of chronic bronchitis, lobar and bronchopneumonia.
29	Urinary tract infections e.g. cystitis, urethritis, pyelonephritis
30	Skin and soft tissue infections e.g. cellulitis, animal bites.

- 31 Dental infections e.g. severe dental abscess with spreading cellulitis.
- 32 Susceptibility to AUGMENTIN will vary with geography and time (see Pharmacological
- 33 Properties, Pharmacodynamics for further information). Local susceptibility data should
- 34 be consulted where available, and microbiological sampling and susceptibility testing
- 35 performed where necessary.
- 36 Mixed infections caused by amoxicillin-susceptible organisms in conjunction with
- 37 AUGMENTIN susceptible beta-lactamase-producing organisms may be treated with
- 38 AUGMENTIN. These infections should not require the addition of another antibiotic
- resistant to beta-lactamases.

40 Dosage and Administration

- Dosage depends on the age, weight and renal function of the patient and the severity of
- 42 the infection.
- Dosages are expressed throughout in terms of amoxicillin/clavulanate content except
- 44 when doses are stated in terms of an individual component.
- To minimise potential gastrointestinal intolerance, administer at the start of a meal. The
- absorption of *AUGMENTIN* is optimised when taken at the start of a meal.
- 47 Treatment should not exceed 14 days without review.
- 48 Therapy can be started parenterally and continued with an oral preparation.
- 49 AUGMENTIN bottle presentations for suspension may be supplied with a plastic dosing
- device. For preparation of the suspensions see *Instructions for Use/Handling*.
- 51 The usual recommended daily dosage is:
- Lower dose: 25/3.6 to 45/6.4 mg/kg/day in two divided doses for mild to moderate
- 53 infections (upper respiratory tract infections e.g. recurrent tonsillitis, lower
- respiratory infections and skin and soft tissue infections).
- *Higher dose*: 45/6.4 to 70/10 mg/kg/day in two divided doses for the treatment of
- more serious infections (upper respiratory tract infections e.g. otitis media and
- 57 sinusitis, lower respiratory tract infections e.g. bronchopneumonia and urinary tract
- 58 infections).
- No clinical data are available on doses above 45/6.4 mg/kg/day in children under 2 years.
- There are no clinical data for AUGMENTIN BD S and SF to make dosage
- 61 recommendations for children under 2 months old.
- 62 The tables below give dosage guidance for children.

Children 2 years and over

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25/3.6	2 - 6 years	5.0 ml AUGMENTIN BD S twice daily or
mg/kg/day	(13 - 21 kg)	2.5 ml AUGMENTIN BD SF twice daily.
	7 - 12 years	10.0 ml AUGMENTIN BD S twice daily or
	(22 - 40 kg)	5.0 ml AUGMENTIN BD SF twice daily
45/6.4	2 - 6 years	10.0 ml AUGMENTIN BD S twice daily or
mg/kg/day	(13 - 21 kg)	5.0 ml AUGMENTIN BD SF twice daily
	7 - 12 years	10.0 ml AUGMENTIN BD SF twice daily.

65 66

Children aged 2 months to under 2 years

AUGMENTIN BD SF suspension 457 mg/5 mL				
Body Weight (kg)	Lower dose at 25/3.6 mg/kg/day (mL every 12 hours)	Higher dose at 45/6.4 mg/kg/day (mL every 12 hours)		
2	0.3	0.6		
3	0.5	0.8		
4	0.6	1.1		
5	0.8	1.4		
6	0.9	1.7		
7	1.1	2.0		
8	1.3	2.3		
9	1.4	2.5		
10	1.6	2.8		
11	1.7	3.1		
12	1.9	3.4		
13	2.0	3.7		
14	2.2	3.9		
15	2.3	4.2		

67 **Renal Impairment**

- No adjustment in dose is required in patients with creatinine clearance greater than
- 69 30 mL/min.

- 70 AUGMENTIN BD S and AUGMENTIN BD SF are not recommended in patients with a
- 71 creatinine clearance of less than 30 mL/min.

72 **Hepatic Impairment**

- 73 Dose with caution; monitor hepatic function at regular intervals. There is, as yet,
- insufficient evidence on which to base a dosage recommendation.

Contraindications

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- 76 AUGMENTIN is contraindicated in patients with a history of hypersensitivity to beta-
- 77 lactams, e.g. penicillins and cephalosporins.
- 78 AUGMENTIN is contraindicated in patients with a previous history of AUGMENTIN-
- 79 associated jaundice/hepatic dysfunction.

Warnings and Precautions

- 81 Before initiating therapy with *AUGMENTIN*, careful enquiry should be made concerning
- previous hypersensitivity reactions to penicillins, cephalosporins or other allergens.
- 83 Serious and occasionally fatal hypersensitivity reactions (including anaphylactoid and
- severe cutaneous adverse reactions) have been reported in patients on penicillin therapy.
- 85 These reactions are more likely to occur in individuals with a history of penicillin
- 86 hypersensitivity (see *Contraindications*). If an allergic reaction occurs, *AUGMENTIN*
- 87 therapy must be discontinued and appropriate alternative therapy instituted. Serious
- anaphylactic reactions require immediate emergency treatment with adrenaline. Oxygen,
- 89 intravenous (i.v.) steroids and airway management (including intubation) may also be
- 90 required.
- 91 AUGMENTIN should be avoided if infectious mononucleosis is suspected since the
- 92 occurrence of a morbilliform rash has been associated with this condition following the
- 93 use of amoxicillin.
- Prolonged use may also occasionally result in overgrowth of non-susceptible organisms.
- 95 Pseudomembranous colitis has been reported with the use of antibiotics and may range in
- severity from mild to life-threatening. Therefore, it is important to consider its diagnosis
- 97 in patients who develop diarrhoea during or after antibiotic use. If prolonged or
- 98 significant diarrhoea occurs or the patient experiences abdominal cramps, treatment
- should be discontinued immediately and the patient investigated further.
- Abnormal prolongation of prothrombin time (increased INR) has been reported rarely in
- patients receiving AUGMENTIN and oral anticoagulants. Appropriate monitoring should
- be undertaken when anticoagulants are prescribed concurrently. Adjustments in the dose
- of oral anticoagulants may be necessary to maintain the desired level of anticoagulation.

104 105 106	Changes in liver function tests have been observed in some patients receiving <i>AUGMENTIN</i> . The clinical significance of these changes is uncertain but <i>AUGMENTIN</i> should be used with caution in patients with evidence of hepatic dysfunction.
107 108 109	Cholestatic jaundice, which may be severe, but is usually reversible, has been reported rarely. Signs and symptoms may not become apparent for up to six weeks after treatment has ceased.
110 111	In patients with renal impairment <i>AUGMENTIN</i> BD S bd L and L BD SF are not recommended.
112 113 114 115	In patients with reduced urine output, crystalluria has been observed very rarely, predominantly with parenteral therapy. During the administration of high doses of amoxicillin, it is advisable to maintain adequate fluid intake and urinary output in order to reduce the possibility of amoxicillin crystalluria (see <i>Overdose</i>).
116 117	AUGMENTIN BD S and BD SF suspensions contain 12.5 mg aspartame per 5 mL dose and therefore care should be taken in patients with phenylketonuria.
118	Interactions
119 120 121	Concomitant use of probenecid is not recommended. Probenecid decreases the renal tubular secretion of amoxicillin. Concomitant use with <i>AUGMENTIN</i> may result in increased and prolonged blood levels of amoxicillin but not of clavulanate.
122 123 124	Concomitant use of allopurinol during treatment with amoxicillin can increase the likelihood of allergic skin reactions. There are no data on the concomitant use of <i>AUGMENTIN</i> and allopurinol.
125 126	In common with other antibiotics, <i>AUGMENTIN</i> may affect the gut flora, leading to lower oestrogen reabsorption and reduced efficacy of combined oral contraceptives.
127 128 129 130	In the literature there are rare cases of increased international normalised ratio in patients maintained on acenocoumarol or warfarin and prescribed a course of amoxicillin. If coadministration is necessary, the prothrombin time or international normalised ratio should be carefully monitored with the addition or withdrawal of <i>AUGMENTIN</i> .
131 132 133 134	In patients receiving mycophenolate mofetil, reduction in pre-dose concentration of the active metabolite mycophenolic acid of approximately 50% has been reported following commencement of oral amoxicillin plus clavulanic acid. The change in pre-dose level may not accurately represent changes in overall MPA exposure.
135	Pregnancy and Lactation
136 137 138 139 140	Reproduction studies in animals (mice and rats) with orally and parenterally administered <i>AUGMENTIN</i> have shown no teratogenic effects. In a single study in women with preterm, premature rupture of the foetal membrane (pPROM), it was reported that prophylactic treatment with <i>AUGMENTIN</i> may be associated with an increased risk of necrotising enterocolitis in neonates. As with all medicines, use should be avoided in

141 142	pregnancy, especiphysician.	cially during the first trimester, unless considered essential by the
143 144 145	the risk of sensiti	ay be administered during the period of lactation. With the exception of isation, associated with the excretion of trace quantities in breast milk, mental effects for the infant.
146	Effects on Ab	oility to Drive and Use Machines
147	Adverse effects of	on the ability to drive or operate machinery have not been observed.
148	Adverse Rea	ctions
149 150 151 152	rare undesirable those occurring a	clinical trials were used to determine the frequency of very common to effects. The frequencies assigned to all other undesirable effects (i.e., at < 1/10,000) were mainly determined using post-marketing data and any rate rather than a true frequency.
153	The following co	onvention has been used for the classification of frequency:
154	very com	$mon \ge 1/10$
155	common	$\geq 1/100 \text{ to} < 1/10$
156	uncommo	on $\geq 1/1000$ to $< 1/100$
157	rare $\geq 1/1$	0,000 to < 1/1000
158	very rare	< 1/10,000.
159	Infections and i	nfestations
160	Common	Mucocutaneous candidiasis
161	Blood and lymp	phatic system disorders
162	Rare	Reversible leucopenia (including neutropenia) and thrombocytopenia
163 164	Very rare	Reversible agranulocytosis and haemolytic anaemia. Prolongation of bleeding time and prothrombin time.
165	Immune system	disorders
166 167	Very rare	Angioneurotic oedema, anaphylaxis, serum sickness-like syndrome, hypersensitivity vasculitis

168	Nervous system	disorders
169	Uncommon	Dizziness, headache
170 171 172	Very rare	Reversible hyperactivity, <i>aseptic meningitis</i> , convulsions. Convulsions may occur in patients with impaired renal function or in those receiving high doses.
173	Gastrointestina	l disorders
174	Adults	
175	Very common	Diarrhoea
176	Common	Nausea, vomiting
177	Children	
178	Common	Diarrhoea, nausea, vomiting
179	All populations	
180 181		often associated with higher oral dosages. If gastrointestinal reactions are by be reduced by taking <i>AUGMENTIN</i> at the start of a meal.
182	Uncommon	Indigestion
183 184	Very rare	Antibiotic-associated colitis (including pseudomembranous colitis and haemorrhagic colitis – see <i>Warnings and Precautions</i>)
185		Black hairy tongue
186 187 188		Superficial tooth discolouration has been reported very rarely in children. Good oral hygiene may help to prevent tooth discolouration as it can usually be removed by brushing.
189	Hepatobiliary d	lisorders
190 191 192	Uncommon	A moderate rise in AST and/or ALT has been noted in patients treated with beta-lactam class antibiotics, but the significance of these findings is unknown.
193 194	Very Rare	Hepatitis and cholestatic jaundice. These events have been noted with other penicillins and cephalosporins.
195 196 197	•	have been reported predominantly in males and elderly patients and may th prolonged treatment. These events have been very rarely reported in

198 199 200 201 202 203	may not become usually reversible deaths have been	toms usually occur during or shortly after treatment but in some cases apparent until several weeks after treatment has ceased. These are le. Hepatic events may be severe and in extremely rare circumstances, in reported. These have almost always occurred in patients with serious use or taking concomitant medications known to have the potential for
204	Skin and subcu	taneous tissue disorders
205	Uncommon	Skin rash, pruritus, urticaria
206	Rare	Erythema multiforme
207 208 209 210	Very rare	Stevens-Johnson syndrome, toxic epidermal necrolysis, bullous exfoliative-dermatitis, acute generalised exanthemous pustulosis (AGEP), and drug reaction with eosinophilia and systemic symptoms (DRESS)
211	If any hypersens	sitivity dermatitis reaction occurs, treatment should be discontinued.
212	Renal and urin	ary disorders
213	Very rare	Interstitial nephritis, crystalluria (see Overdose)
214	Overdose	
215 216 217		symptoms and disturbance of the fluid and electrolyte balances may be ntestinal symptoms may be treated symptomatically with attention to the balance.
218 219	Amoxicillin crys Warnings and P	stalluria, in some cases leading to renal failure, has been observed (see recautions).
220	AUGMENTIN c	an be removed from the circulation by haemodialysis.
221	PHARMACO	DLOGICAL PROPERTIES
222	Pharmacody	namics
223 224 225 226 227	antibiotic before anticipates this or rendering the or	any antibiotics is caused by bacterial enzymes which destroy the it can act on the pathogen. The clavulanate in <i>AUGMENTIN</i> suspension defence mechanism by blocking the beta-lactamase enzymes, thus ganisms sensitive to amoxicillin's rapid bactericidal effect at eadily attainable in the body.
228 229 230	amoxicillin as A	tself has little antibacterial activity; however, in association with <i>UGMENTIN</i> it produces an antibiotic agent of broad spectrum with wide ospital and general practice.

In the list below, organisms are categorised according to their *in vitro* susceptibility to *AUGMENTIN*.

In vitro susceptibility of micro-organisms to AUGMENTIN

Where clinical efficacy of Augmentin has been demonstrated in clinical trials this is indicated with an asterisk (*).

Organisms that do not produce beta-lactamase are identified (with †). If an isolate is susceptible to amoxicillin, it can be considered susceptible to *AUGMENTIN*.

Commonly susceptible species

Gram-positive aerobes:

Bacillius anthracis

Enterococcus faecalis

Listeria monocytogenes

Nocardia asteroides

Streptococcus pyogenes*†

Streptococcus agalactiae*†

Streptococcus spp. (other beta-hemolytic)*†

Staphylococcus aureus (methicillin susceptible)*

Staphylococcus saprophyticus (methicillin susceptible)

Coagulase negative staphylococcus (methicillin susceptible)

Gram-negative aerobes:

Bordetella pertussis

Haemophilus influenzae*

Haemophilus parainfluenzae

Helicobacter pylori

Moraxella catarrhalis*

Neisseria gonorrhoeae

Pasteurella multocida

Vibrio cholerae

Other:

Borrelia burgdorferi

Leptospira ictterohaemorrhagiae

Treponema pallidum

Gram positive anaerobes:
Clostridium spp.
Peptococcus niger
Peptostreptococcus magnus
Peptostreptococcus micros
Peptostreptococcus spp.
Gram-negative anaerobes:
Bacteroides fragilis
Bacteroides spp.
Capnocytophaga spp.
Eikenella corrodens
Fusobacterium nucleatum
Fusobacterium spp.
Porphyromonas spp.
Prevotella spp.
Species for which acquired resistance may be a problem
Gram-negative aerobes:
Escherichia coli*
Klebsiella oxytoca
Klebsiella pneumoniae*
Klebsiella spp.
Proteus mirabilis
Proteus vulgaris
Proteus spp.
Salmonella spp.
Shigella spp.
Gram-positive aerobes:
Corynebacterium spp.
Enterococcus faecium
Streptococcus pneumoniae* [†]
Viridans group streptococcus

	Inherently resistant organisms
	Gram-negative aerobes:
	Acinetobacter spp.
	Citrobacter freundii
	Enterobacter spp.
	Hafnia alvei
	Legionella pneumophila
	Morganella morganii
	Providencia spp.
	Pseudomonas spp.
	Serratia spp.
	Stenotrophomas maltophilia
	Yersinia enterolitica
	Others:
	Chlamydia pneumoniae
	Chlamydia psittaci
	Chlamydia spp.
	Coxiella burnetti
	Mycoplasma spp.
233 234 235 236	Infections caused by amoxicillin-susceptible organisms are amenable to <i>AUGMENTIN</i> treatment due to its amoxicillin content. Mixed infections caused by amoxicillin-susceptible organisms in conjunction with <i>AUGMENTIN</i> -susceptible beta-lactamase producing organisms may therefore be treated with <i>AUGMENTIN</i> .
237	Pharmacokinetics
238	Absorption
239 240 241 242	The two components of <i>AUGMENTIN BD S</i> and <i>AUGMENTIN BD SF</i> , amoxicillin and clavulanate, are each fully dissociated in aqueous solution at physiological pH. Both components are rapidly and well absorbed by the oral route of administration. Absorption of <i>AUGMENTIN</i> is optimised when taken at the start of a meal.
243 244 245 246 247	The mean AUC values for amoxicillin are essentially the same following twice a day dosing with the <i>AUGMENTIN</i> 875/125 mg tablet or three times a day dosing with the <i>AUGMENTIN</i> 500/125 mg tablet, in adults. No differences between the 875 mg twice daily and 500 mg three times daily dosing regimes are seen when comparing the amoxicillin $T_{1/2}$, or C_{max} after normalisation for the different doses of amoxicillin

248249		ed. Similarly, no di er appropriate dose			for the clay	vulanate T _{1/2}	2, C _{max} or A	UC
250 251 252 253 254 255 256	on the pha 875/125 m on the pha mean value AUGMEN	of dosing of AUGM rmacokinetics of ar ag tablet, the time or rmacokinetics of cles and smallest inter TIN at the start of a art of a meal.	noxicilling f dosing tavulanate er-subject	n in adults. relative to e. For clav variabiliti	In a study ingestion of ulanate Al es were ac	of the AUG of a meal had JC and C _{ma} chieved by a	EMENTIN d a marked ef X, the highest dministering	fect
257 258 259		Cmax, Tmax, T1/2 an 875 mg/125 mg neal.					_	
260	Me	an Pharmacokineti	c Parame	eters				
		rug dministration	Dose (mg)	Cmax (mg/L)	Tmax* (hours)	AUC (mg.h/L)	T1/2 (hours)	
	Al	<i>UGMENTIN</i> 1 g						
	Aı	moxicillin	875	12.4	1.5	29.9	1.36	
		avulanate	125	3.3	1.3	6.88	0.92	
261	*M	ledian values						
262 263		in serum concentrate by the oral administ						
264	Distributi	on						
265 266 267		nacokinetics of the te and amoxicillin h						
268	Doubling the dosage of AUGMENTIN approximately doubles the serum levels achieved.							
269	Pre-clini	cal Safety Data	1					
270	No further	information of rele	evance.					
271	PHARM	ACEUTICAL F	PARTIC	CULARS	;			
272	List of E	xcipients						
273 274 275	methylcell	TIN dry powder for ulose, colloidal sili aspberry, orange "l	ca, succi	nic acid, si	licon diox	ide, aspartar		

Incompatibilities

None known.

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278 Shelf Life

The expiry date is indicated on the packaging.

280 Special Precautions for Storage

- Do not take after the expiry date shown on the pack.
- 282 Store in a dry place at or below 30 °C in the original packaging to protect from moisture.
- 283 Refer to pack for storage temperature.
- Once reconstituted, the suspension must be stored in a refrigerator (2°C to 8°C) and used
- within 7 days. Do not freeze. (see also *Instructions for Use/Handling*).

Nature and Contents of Container

- 287 Clear glass bottles containing powder for reconstitution. Bottles may be supplied with
- either an aluminium screw cap with a ring seal or a plastic child-resistant cap with a
- removable foil-backed seal on the bottle. Fill-lines are indicated on the bottle label.
- 290 Bottles may be supplied with a plastic dosing device.

Instructions for Use/Handling

- 292 For bottles with aluminium screw caps, check the cap ring seal is intact before using.
- 293 Alternatively, for bottles with a plastic child-resistant cap, check the foil-backed bottle
- seal is intact before using.
- At time of use, the dry powder should be reconstituted to form an oral suspension, as
- 296 detailed below:
- Invert and shake bottle to loosen powder.
- Add volume of water (indicated below). Invert and shake well.
- Alternatively, fill the bottle with water to just below the mark on bottle label.
- Invert and shake well, then top up with water to the mark. Invert and shake again.
- Allow to stand for 5 minutes to ensure full dispersion.
- Shake well before taking each dose.

AUGMENTIN BD S suspension 228 mg/5 mL				
Fill Weight Volume of water to be added to		Final volume of reconstituted oral		
(g) reconstitute (mL)		suspension (mL)		
7.7	64	70		
15.4	128	140		

AUGMENTIN BD SF suspension 457 mg/5 mL				
Fill Weight Volume of water to be added to reconstitute (mL)		Final volume of reconstituted or suspension (mL)		
6.3	31	35		
12.6	62	70		
25.2	124	140		

304 305 A plastic dosing device may be supplied with the pack which can be used to measure the 306 dose accurately. 307 Discard any unused suspension after 7 days. 308 309 Any unused medicinal product or waste material should be disposed of in accordance with local requirements. 310 Not all presentations are available in every country. 311 312 313 Name and address of the holder of the certificate of registration 314 GlaxoSmithKline South Africa (Pty) Ltd 315 57 Sloane Street 316 Bryanston, 2021 317 South Africa 318 319 Manufacturer 320 Manufacturer: 321 Glaxo Wellcome Production, ZI de la Peyenniere, 53100 Mayenne Cedex, France 322 323 324 **Registration details** 325 Botswana: AUGMENTIN BD SF – Reg No BOT1502714 S2 326

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328 Malawi:

329 AUGMENTIN BD S – Reg No PMPB/PL270/84 POM

AUGMENTIN BD SF – Reg No PMPB/PL270/183 POM 330

331

332	Namibia:
333	AUGMENTIN BD S – Reg No 04/20.1.2/1735 NS2
334	AUGMENTIN BD SF – Reg No 04/20.1.2/1736 NS2
335	
336	Zambia:
337	AUGMENTIN BD S – Reg No 179/009 POM
338	AUGMENTIN BD SF – Reg No 179/046 POM
339	
340	Version number: GDS26/IPI14
341	Date of issue: 13 June 2019
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