The state of the second

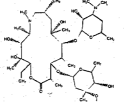
## AZITHROMYCIN FOR ORAL SUSPENSION, USP

To reduce the development of drug-resistant bacteria and maintain the effectiveness of aziltromycin and other antibacterial drugs, aziltromycin should be used only to treat or prevent intections that are proven or strongly suspected to be caused by bacteria.

Strongly suspected to be caused by bacteria.

DESCRIPTION
Azithromycin for oral suspension, USP contain the active inprecient azithromycin, an azailde, a subclass of
macroided ambitiotics. for oral administration. Azithromycin has the chemical name
(28.35.48.58.88.108, 118.11.25.135.146):13-1(12.6-dideoxy-2-C-mathyt-a

Azithromycin has the following structural formula: Azithromycin has the following structural formula: Azithromycin, as the monohydrate, is a white crystalline powder with a molecular formula of CapitryACQ-pt-Q and a molecular weight of 767 0. Azithromycin for oral suspension, SSP is supplied as azithromycin monohydrate powder equivalent to 300 mg, 500 mg, 900 mg or 1200 mg azithromycin expension or per bottle and the following inactive ingredients: cololoidal silicon closide, Protorygropy Cellulose, sucrose, tribasic sodium phosphate, anhydrous, santhan gum, beannal flavor, cherry flavor, vanilla flavor and FD&C Red No. 40. After constitution, each 5 mt. of suspension contains 100 mg or 200 mg of azithromycin.



# CLINICAL PHARMACOLOGY

CLIMICAL PRAIMMEDULA.

Pharmacokineth mean (SD) pharmacokinetic parameters were  $\text{AUC}_{0,T^2} = 4.3$  (1.2) mcg/h/ml;  $C_{\text{max}} = 0.5$  (0.2) mg/ml;  $C_{\text{max}} = 0.5$  (0.2) mcg/ml;  $C_{\text{max}} =$ 

Pharmacokinetic Parameters (Mean)	Total n=12	Davi 6
C <sub>max</sub> (mcg/mL)	Day 1 0.41	<u>Day 5</u> 0.24
Tensy (h)	2.5	3.2
AUC <sub>0-24</sub> (mcg-h/mL)	2.6	2.1
C <sub>min</sub> (mcg/mL)	0.05	0.05
Urinary Excret. (% dose)	4.5	6.5

Azitnromycin 250 mg tablets are bioequivalent to 250 mg capsules mg capsules are no longer commercially available.

ing a two-way crossover study, 12 adult healthy volunteers (6 males, 6 females) received 1,500 mg of azithromycin administered in single daily doses over either 5 days (two 250 mg tablets on day 1, followed



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by one 250 mg lablet on days 2-5) or 3 days 1500 mg per day for days 1-3). Due to limited serum samples on day 2 (3-day retimen) and days 2-4 (5-day retimen), the serum concentration-time profile of each subject was fit to a 3-compartment model and the AUCo\_ for the fitted concentration profile was comparable between the 5-day and 3-day retimens.

	3-Day l	Regimen	5-Day I	Regimen
Pharmacokinetic Parameter	Day 1	Day 3	Day 1	Day 5
[mean (SD)] C <sub>mb</sub> (serum, mcg/mL)	0.44 (0.22)	0.54 (0.25)	0.43 (0.20)	0.24 (0.06)
Serum AUC <sub>0</sub> (mcg-hr/mL) Serum T <sub>1/2</sub>	17.4 71.	(6.2) 8 hr		(3.1)* 1.9 hr

Median authrougic exposure (AUC<sub>0-208</sub>) in mononuclear (ANN) and polymorphonuclear (PMN) leukocytes following either the 5-day or 3-day regimen was more than a 1000-100I and 800-1010 greater than in serum, respectively, Administration of the same total does with either the 5-day or 3-day regimen may be expected to provide comparable concentrations of authromycin within NN and PMN leukocytes.

Two authromycin 250 mp tablets are bloeduivalent to a single 500 mg tablet.

I wo azimromycin 250 mg labeles are bioequivalent to a simple 500 mg labele.

Abbargtion

The absolute bioavaliability of azithromycin 250 mg capsules is 38%.

The absolute bioavaliability of azithromycin 250 mg capsules is 38%.

In a two-way crossover study in which 12 healthly subjects received a single 500 mg dose of azithromycin (two 250 mg tablets) with or without a high lat meal, food was shown to increase  $C_{ma_k}$  by 23% but had no effect on AUC.

When azithromycin suspension was administered with food to 28 adult healthy male subjects.  $C_{ma_k}$  increased by 55% and AUC was onchanged.

The AUC of azithromycin was unaffected by co-administration of an antical containing aluminum and magnesium hydroide with azithromycin capsules: however, the  $C_{ma_k}$  was reduced by 24%. Administration of cimelidine (800 mg) two hours prior to azithromycin had no effect on azithromycin absorption.

Distribution
The serum protein binding of azithromyoin is variable in the concentration range approximating human exposure, decreasing from \$15 at 0.02 mag/mil. to 7% at 2 mag/mil.
The serum protein binding of azithromyoin to 7% at 2 mag/mil.
The serum protein binding of \$1.00 mag/mil. to 7% at 2 mag/mil.
The serum protein binding of \$1.00 mag/mil. to 7% at 2 mag/mil.
The protein of \$1.00 mag/

AZITHROMYCIN CONCENTRATIONS FOLLOWING A 500 mg DOSE (TWO 250 mg CAPSULES) IN ADULTS

TISSUE OR FLUID	TIME AFTER DOSE (h)	TISSUE OR FLUID CONCENTRATION (mcg/g or mcg/mL)	CORRESPONDING PLASMA OR SERUM LEVEL (mcg/mL)	TISSUE (FLUID) PLASMA (SERUM) RATIO			
SKIN	72-96	0.4	0.012	35			
LUNG	72-96	4.0	0.012	>100			
SPUTUM"	2-4	1.0	0.64	2			
SPUTUM"	10-12	2.9	0.1	30			
TONSIL***	9-18	4.5	0.03	>100			
TONSIL***	180	0.9	0.006	>100			
CERVIX	19	2.8	0.04	70			

CERVIX\*\*\*

19
2.8
0.04
70

1 Azithromych tissue concentrations were originally determined using 250 mg capsules.
Sample was obtained 2-4 hours after the left of the second of the left of the left of the second of the left of the l

Elimination
Plasma concentrations of azithromycin following single 500 mg oral and i.v. doses declined in a polyphasic patient with a mean apparent plasma clearance of 630 mU/min and terminal elimination half-life is flought to be due to extensive uptake and subsequent release of drug from tissues.

iron issues.

Biliary excretion of azilitromycin, predominantly as unchanged drug, is a major route of elimination. Over the course of a week, approximately 6% of the administered dose appears as unchanged drug in urine.

course or a week, approximately one of the administrated oose appears as unchanged drug in unine. Special Populations Renal Insufficiency Azilhomycin pharmacokinetics were investigated in 42 adults (21 to 85 years of age) with varying degrees of renal impairment. Following the oral administration of a single 1,000 mg dose of azilhomycin, meat  $G_{m_1}$  and  $G_{M_2}$  and  $G_$ 

dosage adjustment is recommended based on gender. 
Berlatine Pallems
When studied in healthy elderly subjects aged 65 to 85 years, the pharmacokinelic parameters of adithromycin in elderly men were similar to those in young adults; however, in elderly women, although higher peak concentrations (increased by 30 to 50%) were observed, no significant accumulation occurred. Pediatric Pallets:
In two clinical studies, arithromycin for oral suspension was dosed at 10 mg/kg on day 1, followed by 5 mg/kg on days 2 through 5 to two groups of pediatric palients (apped 1-5 years and 5-15 years, respectively). 
The mean pharmacokinelic parameters on day 5 were C<sub>lonic</sub>0,216 mg/ml. T<sub>lonic</sub>1.9 hours, and AUC<sub>Gu</sub>2.3.028 mg/ml/ml. for the 5- to 5-year-old group, and were C<sub>lonic</sub>0.333 mg/ml., T<sub>lonic</sub>2.4 hours, and AUC<sub>Gu</sub>2.3.108 mg/ml/ml. for the 5- to 15-year-old group.
Two clinical studies were conducted in 65 pediatric palents aged 3-16 years to determine the pharmacokinelics and safety of azithromycin for oral suspension. Arithromycin was administered following a low-lat breakfast.

The lists study consisted of 35 pediatric palents treated with 20 mg/kg/day (maximum daily dose 500 mg) for 3 days of hom 34 patients were evaluated for pharmacokinelics.

placing interests of the property of the prope

3-Day Regimen (20 mg/kg x 3 days) 11 1.1 (0.4) 2.7 (1.9) 7.9 (2.9) 5-Day Regimen (12 mg/kg x 5 days) Pharmacokinetic Parameter [mean (SD)] n C<sub>max</sub> (mcg/mL) T<sub>max</sub> (hr) AUC<sub>0-24</sub>(mcg-hr/mL)

AÜC<sub>0-2</sub> (mcp-h/mL) 7.9 [2.5] 3.9 (1.9]

The similarity of the overall exposure (AUC<sub>0--</sub>) between the 3-day and 5-day regimens in pediatric patients is unknown.

The similarity of the overall exposure (AUC<sub>0--</sub>) between the 3-day and 5-day regimens in pediatric patients is unknown.

Single dose pharmacokinetics in pediatric patients given doses of 30 mg/kg have not been studied, (See DOSAGE AND ADMHISTRATION.)

Drug-Drug Interactions studies were performed with azikhromycin and other drugs likely to be co-administered. The effects of co-administration of azikhromycin on the pharmacokinetics of other drugs are shown in Table 1 and the effect of other drugs on the pharmacokinetics of azikhromycin in Table 2.

Co-administration of azikhromycin at interapetric doses had a modest effect on the pharmacokinetics of azikhromycin, Netficand significantly increased the C<sub>max</sub> and AUC of azikhromycin. No dosage adjustment of azikhromycin, Netficand significantly increased the C<sub>max</sub> and AUC of azikhromycin. No dosage adjustment of azikhromycin is recommended when administered with drugs listed in Table 2. (see PRECAUTIONS-Ong lateractions.)

Table 1. Drug Interactions: Pharmacokinetic Parameters for Co-administered Drugs in the

Table 1. Drug Interactions: Pharmacokinetic Parameters for Co-administered Drugs in the

Co-administered Drug	Dose of Co-administered Drug	Dose of Azithromycin	П	Ratio (with/without of Co-administ Pharmacokinet (90% CI); No	Hered Drug
		Mean C <sub>max</sub>	Mean AUC		
Atorvaslatin	10 mg/day x 8 days	500 mg/day PO on days 6-8	12	0.83 (0.63 to 1.08)	1.01 (0.81 to 1.25)
Carbamazepine	200 mg/day x 2 days, then 200 mg BIO x 18 days	500 mg/day PO for days 16-18	7	0.97 (0.88 to 1.06)	0.96 (0.88 to 1.06)
Cetirizine	20 mg/day x 11 days	500 mg PO on day 7, then 250 mg/day on days 8-11	14	1.03 (0.93 to 1.14)	1.02 (0.92 to 1.13)
Didanosine	200 mg PO BID x 21 days	1,200 mg/day PO on days 8-21	6	1.44 (0.85 to 2.43)	1,14 (0.83 to 1.57)
Elavirenz	400 mg/day x 7 days	600 mg PO on day 7	14	1.04*	0.95*
Fluconazole	200 mg PO single dose	1,200 mg PO single dase	18	1.04 (0.98 to 1.11)	1.01 (0.97 to 1.05)
Indinavir	800 mg TID x 5 days	1,200 mg PO on day 5	18	0.96 (0.86 to 1.08)	0.90 (0.81 to 1.00)
Midazolam	15 mg PO on day 3	500 mg/day PO x 3 days	12	1.27 (0.89 to 1.81)	1.26 (1.01 to 1.56)
Nellinavir	750 mg TID x 11	1.200 mg PO on day	14	0.90	0.85 (0.78 to (1.93)



Ritabutin	300 mg/day x 10 days	500 mg PO on day 1, then 250 mg/day on days 2-10	6	See tootnote below	NA .
Sildenafil	100 mg on days 1 and 4	500 mg/day PO x 3 days	12	1.16 (0.86 to 1.57)	0.92 (0.75 to 1.12)
Theophylline	4 mg/kg IV on days 1, 11, 25	500 mg PO on day 7, then 250 mg/day on days 8-11	10	1.19 (1.02 to 1.40)	1.02 (0.86 to 1.22)
Theophylline	300 mg PO BID x 15 days	500 mg PO on day 6, then 250 mg/day on days 7-10	8	1.09 (0.92 to 1.29)	1.06 (0.89 to 1.31)
Triazolam	0.125 mg on day 2	500 mg PO on day 1, then 250 mg/day on day 2	12	1.06	1.02*
Trimethoprim/ Sullamethoxazole	160 mg/800 mg/day PO x 7 days	1,200 mg PO on day day 7	12	0.85 (0.75 to 0.97) (0.90 (0.78 to 1.03)	0.87 (0.80 to 0.95) /0.96 (0.88 to 1.03)
Zidovudine	500 mg/day PO x 21 days	600 mg/day PO x 14 days	5	1.12 (0.42 to 3.02)	0.94 (0.52 to 1.70)
Zidovudine	500 mg/day PO x 21 days	1,200 mg/day PO x 14 days	4	1.31 (0.43 to 3.97)	1.30 (0.69 to 2.43)

NA - Not Available

- 99% Confidence interval not reported

Mean rifabutin concentrations one-half day after the last dose of rifabutin were 60 ng/mL when coadministered with arithromycin and 71 ng/mL when co-administered with placebo.

Pharmacokinetic Parameters for Arithromycin in the Presence of Co-

Table 2. Drug Interactions: Pharmacokinetic Parameters for Arithromycin in the Presence of Co

Co-administered Drug	Dose of Co-administered Drug	Dose of Azithromycin	U	co-administr Azithromycin F Paramete	th/without ered drug) of Pharmacokinetic rs (90% CI); est = 1.00
				Mean C <sub>max</sub>	Mean AUC
Etavirenz	400 mg/day x 7 days	600 mg PO on day 7	14	1.22 (1.04 to 1.42)	0.92
Fluconazole	200 mg PO single dose	1,200 mg PO single dose	18	0.82 (0.66 to 1.02)	1,07 (0.94 to 1.22)
Nelfinavir	750 mg TID x 11 days	1,200 mg PO on day 9	14	2.36 (1.77 to 3.15)	2.12 (1.80 to 2.50)
Rifabutin	300 mg/day x 10 days	500 mg PO on day 1, then 250 mg/day on days 2-10	6	See footnote below	NA .

In days 2-10

MA – Not available

- 90% Conlidence interval not reported

Mean azithromycin concentrations one day after the last dose was 53 ng/mL, when coadministered with 300 mg daily rilabutin and 49 ng/mL when coadministered with placebo.

Microbiology: Azithromycin acts by binding to the 505 ribosonnal subunit of susceptible microorganisms and, thus, interfering with microbial protein synthesis. Nucleic acid synthesis is not affected. Azithromycin actonectirates in placopyetes and tibroblasts as demonstrated by in virto includation techniques. Using such methodology, the ratio of intracellular to extracellular concentration was 530 after one hour includation. In vivo studies suggest that concentration in phagocytes may contribute to drug distribution to inflamed tissues. Azithromycin has been shown to be active against most isolates of the following microorganisms, both in virto and in clinical infections as described in the IMDICATIONS AND USAGE section.

wino and in clinical intections as described in the IMDICATIONS AND USAGE section.

Aerobic and textulative gram-positive microorganisms

Streptococcus aguaratise

Streptococcus propones

NOTE: Achircomycin demonstrates cross-resistance with erythromycin-resistant gram-positive strains. Most strains of Entercoccus Repairs and methicillin-resistant staphylococci are resistant to azithromycin. Aerobic and facultative gram-negative microorganisms

Hearmophilis ductery

Hearmophilis influenzae

Moraxella catarrhalis

Neisseria gonorrhoeae "Other" microorganisms Chlarnydia pneumoniae Chlarnydia trachomatis

Chamyols preumoniae

Mycopiasma preumoniae

M

'Other" microorganisms

Complex or weakyfour Unexplosers weakyfour Susceptibility Institut Historia When weaken the Complex of its wice susceptibility test results for antimicrobial drugs used in resident When weaken the resident of the physician as periodic reports which describe the susceptibility profile of nessecomial and community-acquired pathogens. These reports may differ from susceptibility data obtained from outpatient use, but could aid the physician in selecting the most effective antimicrobial.

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(contin or spen / continues).

Biffusion Jechniques:

Countrative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of batteris to antimicrobial compounds. One such standardized procedure<sup>2,5</sup> requires the use of standardized inoculum concentrations. This procedure uses paper disks impregnated with 15-mcg arithnomycin to test the susceptibility of microorganisms to azithnomycin. The disk diffusion interpretive criteria are provided in Table 1.

Table 1. Susceptibility interpretive Criteria for Azithromycin Susceptibility Test Result interpretive Criteria

	Concentrations (meg/mL)		(zone diameters in mm)			
Pathogen	- 1			2.5		
	5		R <sup>2</sup>	S .	: 1	B.
Haemophilus 500.	<b>54</b>			≥ 12		
Staphylococcus aureus	≤ 2	4	≥.8	. ≥18	14-17	. ≤ 13
Streptococci including						
S. pneumoniaeb	≤ 0.5	. 3	≥ 2	≥ 18	14-17	≤ 13
2 The current absence o	I data on	resistant strai	ns preciudes	defining any ca	aleoory other	than "susce

Ine current assence of data on resistant strains precludes defining any category other than "susceptible."
 It strains yield MIC results other than susceptible, they should be submitted to a reference laboratory for further testing.

Susceptibility of streptococci including S. pneumoniae to azithromycin and other macrolides can be predicted by testing enythromycin.

elive criteria have been established for testing Neisseria gonorrhoeae. This species is not usually

tested. A report of "susceptible" indicates that the palhogen is likely to be inhibited if the antimicrobial compound reaches the concentrations usually achievable. A report of "intermediate" indicates that the result should be considered equivocal, and, if the microorganism is not fully susceptible to alternative, clinically lessible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is phyliologically concentrated or in situations where high dosage of drug can be used. This category also provides a buffer zone which prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of "resistant" indicates that the pathogen is not likely to be inhibited in the antimicrobial compound reaches the concentrations usually achievable; other therapy should be

DUALITY CONTROL:

DUALITY DUTINIX:
Standardized susceptibility test procedures require the use of quality control microorganisms to control the technical aspects of the tests procedures. Standard arithmorphic provide should provide the following trange of values noted in Table 2. Quality control microorganisms are specific, starins of prefamiliant minimisc biological properties. OS strains are very stable strains which will give a standard and repeatable susceptibility petition. The specific strains of used to microbiological quality control are not clinically

significant.	Acceptable Quality Control Ranges	las Azithramucia
OC Strain	Minimum inhibitory Concentrations (mcg/mL)	Disk Diffusion (zone diameters in mm)
Haemophilus influenzae ATCC 49247	1.0-4.0	13-21
Staphylococcus aureus ATGC 29213	0.5-2.0	
Staphylococcus aureus ATCC 25923		21-26
Streptococcus pneumoniae ATCC 49619	0.06-0.25	19-25

INDICATIONS AND USAGE
ASthormycin is indicated for the treatment of patients with mild to moderate infections (pneumonia; see
WARNINGS) usaces by susceptible strains of the designated microorganisms in the specific conditions
listed below. As recommended dosages, durations of therapy and applicable patient populations vary among
these intections, please see DSAGE AND ADMINISTRATION to specific dosting occumentalitions.

Acute bacterial exacerbations of chronic obstructive pulmonary disease due to Haemophilus influenzae, Moraxella catarrhalis or Streotococcus pneumoniae.

Moraxella catarrhalis or Streptococcus pneumoniae.

Acute bacterial sinusitis due to Haemophilus influenzae. Moraxella catarrhalis or Streptoc

Acute bacterial sinustits due to \*næmogninas minaseas\*, minaseas\*,

respond to meri timess (including immandesticency of relucional asplenal).

Pharyngillus/nessillist caused by Sierelopoccus progenes as an alternative to first-line therapy in individuals who cannot use irist-line therapy.

NOTE: Peniciain by the intramuscular route is the usual drug of choice in the treatment of Streptococcus progenes meeting and the prophylaxis of rheumatic lever, Authromycin is often effective in the eradication of susceptible statistic of Streptococcus progenes from the associations. Because some strains are existent to arthromycin, susceptibility lests should be performed when patients are treated with authromycin, bate establishing fitting of patients provided in authromycin. Date establishing fitting of patients provided in a susceptible strain of surprise of the strain of the st

estansing efficacy of aziliromyon in subsequent prevention of rheumatic lever are not available. Uncomplicated shis and shis latinuture intections due to Staphyococcus aureus. Streptococcus pyogenes or Streptococcus agalactile. Abscesses usually require surgical drainage. Ureibritis and certifelis due to Chlamydis trachomatis or Neisseris gonomboeae. Geallati ulera disease in men due to Naemophius ducreyl chancord), Due to the small number of women included in clinical trials, the efficacy of azithromycin in the treatment of chancroid in women has not been established.

stabilished.
Azihromycin, at the recommended dose, should not be relied upon to treat syphilis. Antimicrobial agents used in high doses for short periods of time to treat non-gonococcal urethrikis may mask or delay the symptoms of incubaling syphilis. All patients with sexuality-transmitted urethrikis or cervicitis should have seriogic test for syphilis and appropriate autimets for gonomes performed at the time of diagnosis. Appropriate antimicrobial therapy and follow-up tests for these diseases should be initiated if infection is continued.

springic test for syphilis and appropriate and following tests for these diseases should be minimicrobial therapy and following tests for these diseases should be minimicrobial therapy and following tests should be performed before treatment to determine the causative organism and its susceptibility tests should be performed before treatment to determine the causative organism and its susceptibility to azintromycin. Therapy with azintromycin may be initiated before results of organism and its susceptibility to azintromycin. Therapy with azintromycin may should be adjusted to the state of the stat

these tests are known; once the results become available, animicrobial therapy should be adjusted accordingly.

To reduce the development of drug-resistant bacteria and maintain the effectiveness of aziltromycin and other antibacterial drugs, aziltromycin should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptible bacteria. When culture and susceptibility information are variable, they should be considered in selecting or modifying antibilitaterial therapy, in the absence of such data, local epidemiology and susceptibility patterns may conflibility to the empiric selection of therapy. Pediatric Patterns: (See PRECANTIONS—Pediatric Iste and CLINICAL STUDIES IN PEDIATRIC PATTERTS.) Acute otilis media caused by \*haemophibus influenzae, Morazella catarinalis or Stephococcus pneumoniae. (For specific dosage recommendation, see DOSAGE AND ADMINISTRATION.)

Community-acquired pneumonia due to Chiampdia pneumoniae. Haemophibus influenzae, Morazella catarinalis or Stephococcus pneumoniae in plenista appropriate for oral therapy. (For specific dosage recommendation, see DOSAGE AND ADMINISTRATION.)

NOTE: Additionary in should not be used in addiatic pallents with aneumonia who are lutined to be

recommensation, see public and be used in pediatric patients with pneumonia who are judged to be inappropriate for oral therapy because of moderate to severe litness or risk factors such as any of the

patients with cystic fibrosis, patients with noscoemially acquired infections, patients with moore or suspecied bacterenta, patients requiring hospitalization, or patients with significant underlying health problems that may compromise their ability to respond to their illness (including immunodeficiency or functional asplenia). Pharynghis/nonsililits caused by Streptococcus progeness as a alternative birsh-line therapy in ndividuals who cannot use first-line therapy. (For specific dosage recommendation, see DOSAGE AND ADMINISTRATION.)

ADMINISTRATION.)

NDTE: Peniciliin by the intramuscular route is the usual drug of choice in the treatment of Streptococcus progress intection and the prophylaxis of rheumatic lever. Azilhromych is often effective in the eracicular of susceptible strains of Streptococcus progeness inclined in a susceptible strains of Streptococcus progeness from the nasopharym. Because some strains are resistant to azilhromych, susceptibility is studio be performed when patients are resistant as rather with azilhromych as stabilishing efficacy of azilhromycin in subsequent prevention of rheumatic lever are not available. Appropriate outline and susceptibility lests should be performed before treatment to determine the causalive organism and its susceptibility to azilhromycin. Therapy with azilhromycin may be initiated before results of these tests are known; once the results become available, antimicrobial therapy should be adjusted accordingly.

CONTRAINDICATIONS Azithromycin is contrai macrolide antibiolic.

to raindicated in patients with known hypersensitivity to azithromycin, erythromycin or any

WARNINGS

WARNINGS
Serious allergic reactions, including angloedema, anaphylaxis, and dermatologic reactions including Stevens
Johnson Syndrome and toxic epidermal necrobysis have been reported rarely in patients on azithromycin
therapy. Although rare, Isalities have been reported. (See DothTRIMIDICATIONS,) Despite initially
successful symptomatic treatment of the alergic symptoms, when symptomatic therapy was decornitised,
successful symptomatic treatment of the alergic symptoms, when symptomatic therapy was decornitised,
reported by the state of the alergic symptoms and symptomatic threapy was decornitised,
reported by the state of the state these episcoes to the long inside that the drug should be discontinued and appropriate therapy should be instituted.

It an alterior reaction occurs, the drug should be discontinued and appropriate therapy should be instituted.

Physicians should be aware that reappearance of the altergic symptoms may occur when symptomatic therapy is discontinued.

Physicians should be aware that reoppearance of the allergic symptoms intrapy should be instituted, therapy is discontinued. In the treatment of parameters are all the properties of the allergic symptoms and occur when symptomatic therapy is discontinued. Simple properties are all the properties of the allergic symptoms and the continuence of the allergic symptoms and the allergic symptoms are all the allergic symptoms and the allergic symptoms are all the allergic symptoms and the allergic symptoms are all the allergic symptoms and the allergic symptoms are allergic symptoms and allergic symptoms and allergic symptoms are allergic symptoms and allergic symptoms and allergic symptoms are allergic symptoms and allergic symptoms and allergic symptoms are allergic symptoms and allergic symptoms and allergic symptoms are allergic symptoms and allergic symptoms and allergic symptoms and allergic symptoms are allergic symptoms

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RECAUTIONS
General: Because azithromycan is principally eliminated via the liver, caution should be exercised when authromycan is administered to patients with impaired hepatic function. Due to the limited data in subjects with GFR c10 mt\_min. caution should be exercised when prescribing azithromycin in these patients, (see CLINICAL PHAMACOLOGY: Special Populations - Renal lexitaticiency.)
Protonged cardiac repolarization and OT interval, imparting a risk of developing cardiac arrivythmia and lorsades de pointes, have been seen in treatment with other macrolless. A seniar ericle with azithromycin in the absence of a proven or strongly suspected bacterial intection or a prophylactic indication is unlikely to provide benefit to the patient and increases the risk of the development of drug-resistant bacteria. Information for Patients:
Altitromycin or all suspension can be taken with or without food.
Patients should also be cautioned not to take aluminum— and magnesium-containing antacids and azithromycin simultaneously.

Patients should also be cautioned not to take aluminum—and magnesium-containing antacids and arithromycin immultaneously. The patient should be directed to discontinue azithromycin immediately and contact a physician if any signs of an altergic reaction occur. Patients should be counseled that antibacterial drugs including azithromycin should only be used to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When azithromycin is prescribed to treat a bacterial infection step, the common cold. When azithromycin is prescribed to treat a bacterial infection patients should be taken exactly as directed. Skipping doses or not completing the full course of therapy may (1) decrease the effectiveness of the immediate treatment and (2) increase the Reischood that bacteria will develop resistance and will not be treatable by azithromycin or other antibacterial drugs in the tuture.

antibacterial drugs in the future.

Drug Interactions

Co-administration of nelinavir at steady state with a single oral dose of azithromycin resulted in increased azithromycin serum concentrations. Although a dose adjustment of azithromycin is not recommended when administration of nelinavir, close monitoring for known side effects of azithromycin, such as lives enzyme anonramisties and hearing impairment, is warranted. (See ADVERSE REACTIONS). Azithromycin did not affect the prothrombin lime response to a single dose of warfarin. However, prodent medical practice clicates careful monitoring of prothrombin time in all patients treated with azithromycin and warfarin concomitantly. Concurrent use of macrotides and warfarin in clinical practice has been associated with increased anticoagulant effects.

Drug interaction studies were performed with azithromycin and other drugs likely to be co-administered. See CLINICAL PARAMACIOLO-107-lung-Drug Interactions. When used in therapeutic doses, azithromycin lad a modest effect on the pharmacokinetics of althrostatin, carbamazepine, celirizine, didanosine, elevierus, luccinaziole, midianterinosaciole, didovudine, Co-administration, entrevenus and orall-lad are motest effect on the pharmacokinetics of azithromycin. No dosage adjustment of either drug is recommended withen azithromycin is coadministered with any of the above agentic, interactions with the drugs tisted below have not been reported in clinical trais with azithromycin: however, no specific drug interaction studies have been performed to evakuate potential drug-drug interaction. Nonetheless, they have been observed with macrolide products. Until further data are developed reparding interactions with the drugs tisted below have not been reported in clinical trais with azithromycin: however, no specific drug interaction studies have been performed to evakuate potential drug-drug interaction. Nonetheless, they have been observed with macrolide products. Until further data are developed reparding to potamine and p

Laboratory Test Interactions: There are no reported laboratory test interactions.

Carcinogenesia, Mulaganesis, Impairment of Fartillity: Long-term studies in animals have not been performed to evaluate carcinogenic potential. Azibhromych has shown no mulagenic potential in standard laboratory tests; mouse propiona assays, and mouse bone marrow catalogenic assays, and mouse bone marrow. Pregnancy: Teratogenic Ellectis, Pregnancy Category 8: Reproduction studies have been performed in rats and mice at dones up to moderately maternally toxic does conformations (i.e., 200 mg/kg/dg/). These does, based on a retrymit basis, are estimated to be 4 and 2 mess, respectively; the human daily dose of 50 mg in the animal studies, no evaluate on them to the fetus due to astimomycin was found. There are, however, no adequate and well-controlled daties in pregnant women. Because animal reproduction studies are not always predictive of human response, aztimomycin should be tested during pregnancy only if clerity readed. Nursing Mothers: it is not known whether aztimomycin is excreted in human milk. Decause many drugs are excreted in human milk. Decause many drugs are excreted in human milk. Categoria should be excised with aztimomycin sa doministed to a must grow drugs. A properties of the must be added to the pregnancy only if clerity medical. PARIMACOLOGY, INDICATIONS AND USAGE, and DOSAGE AND ADMINISTRATION). Acute Othis Media (total dosage regimen: 30 mg/kg, see DOSAGE AND ADMINISTRATION): Saley and effectiveness in the treatment the treatment and the properties of the properties o

electiveness in the treatment of pediatric patients with office media under 6 months of age have not been electiveness in the treatment of pediatric patients with office media under 6 months of age have not been exceeded in the treatment of pediatric patients with a consistency and effectiveness in the treatment of acute bacterial sinusitis under 6 months of age have not been established. Use of azithcomycin for the treatment of acute bacterial sinusitis in pediatric patients, or a supported by adequate and well-controlled studies in adults, similar pathophysiology of acute sinusitis in adults and pediatric patients, and studies of acute officis media in pediatric patients.

Community-Acquired Preumonia (dosage regiment: 10 mp/kg on Day 1 followed by 5 mp/kg on Days 2-5): Safety and effectiveness in the treatment of pediatric patients.

Community-Acquired Preumonia (dosage regiment: 10 mp/kg on Days 1 followed by 5 mp/kg on Days 2-5): Safety and effectiveness in the treatment of pediatric patients.

Community-Acquired Preumonia (dosage regiment: 10 mp/kg on Bays 1 mp/kg on Days 1 followed by 5 mp/kg on Days 2-5): Safety and effectiveness for preumonia under 6 effectiveness for preumonia with the preumonia with

Azithromycin los oral suspension 200 mg/s ml. contains 8.8 mg of sodium per 6 ml. of constituted solution. ADVERSE REACTIONS
In climaal trials, most of the reported side effects were mild to moderate in swertly and were reversible upon idecontinuation of the drug. Polanishly services side effects of appliederma and cholestatic jaundice were reported at rely. Approximately 0.7% of the patients tadults and pediants patients from the 5-day multiple-dose crinical trials discontinuated actionnych therapy because of treatmen-leaded side effects, in adults person product prate focus of the solution of the

abdominal pain. (See currence. Common treatment-related side effects in adult patients receiving multiple-dose regimens: Overall, the most common treatment-related side effects in adult patients receiving multiple-dose regimens of azithromycin were related to the gastrointestinal system with diarnea/loose stools (4-5%), nauses (3%) and abdominal pain (2-3%) being the most frequently reported. No other treatment-related side effects occurred in patients on the multiple-dose regimens of azithromycin with a frequency greater than 1%. Side effects that occurred with a frequency of 1% or less included the influence.

stools (4-5%), nausea (3%) and abdominal pain (2-3%), being the most Inequenty No other treatment-related side effects occurred in patients on the multiple-doser with a frequency greater than 1%. Side effects that occurred with a frequency of tollowing:
Cardiovascular: Palpitalions, chest pain.
Gestrointestinat: Dysepsia, flatulence, vomiting, melena and cholestatic jaundice.
Gentlourinary; Monita, vaginitis and nephritis.
Nervous System: Dizziness, headache, vertigo and somnolence.

General: Fatique.

Dosage

General: Fatigue.

Altergie: Rash, pruritus, photosensitivity and angioadema.

Single 1-gam dose regimen: Overall, the most common side effects in patients receiving a single-dose regimen of a grain of authoroxycin were related to the gastiotisatimal system and were more frequently reported that the production of authoroxycin was proposed to the patients on the sauge consequent of significant or authoroxycin with a frequency of 1% or greater included distributions of solds (7%), nauses (5%), abdominal pain (5%), comiting (2%), dyspebsis (1%) and vagnistis (1%).

Single 2-grain dose regimen: Overall, the most common side effects in patients receiving a single 2-grain dose of authoroxycin vere related to the gastriotinestimal system. Side effects that occurred in patients in this study with a frequency of 1% or greater included nauses (18%), distributions of control of 1% or greater included nauses (18%), distributions of 10%, vomiting (7%), administ (2%), dyspepsis (1%) and dizziness (1%). The majority of these complaints were mild in nature.

Pediatric Patients:
Single and Multiple-gose regimens: The types of side affects in pediatric patients were comparable to those seen in adults, with different incidence raiss for the dosage regimens recommended in pediatric patients, deute Oithis Medic: For the recommended tolat dosage regimen of 30 mp/dp, the most frequent side effects (2:1%) attributed to treatment were disprime, abdominal pain, vomiting, nausea and rash, (See DOSAGE AND ADMINISTRATION and CLINICAL STUDIES IN "EQUIPATINE CATENTIA").

Dosage		Abdominal			
Regimen	Diarrhea, %	Pain, %	Vomiting, %	Nausea, %	Rash, %
1-day 3-day	4.3% 2.6%	1,4% 1,7%	4.9% 2.3%	1.0% 0.4%	1.0% 0.6%
5-day	1.8%	1.2%	1.1%	0.5%	0.4%

Community-Acquired Pneumonia: For the recommended dosage regimen of 10 mg/kg on Day 1 followed by 5 mg/kg on Days 2-5, the most frequent side effects attributed to treatment were diarrhea/loose stools, abdominal palm, vomitting, nausea and rash.

The incidence is described in the table below:

Regimen	stools, %	Pain, %	Vomiting. %	Nausea. %	Rash, %
5-day	5.8%	1.9%	1,9%	1.9%	1.6%
Pharyngitis/ton	sillitis: For the rec	ommended dosage	regimen of 12 mg	/kg on Days 1-5.	the most frequer

side effects attributed to treatment were diarri The incidence is described in the table below:

Dosage Regimen Diarrhea, % Pain, % 5-day 5.4% 3.4% Vorniting, % Nausea, % Rash, % 0.7% Headache. % 1.1% With any of the treatment regimens, no other treatment-related side effects occurred in pediatric patients treated with azintromycin with a frequency greater than 1%. Side effects that occurred with a frequency of 1% or less included the following.

Cardiovascular: Chest pain. Gastrointestinal: Dyspepsia, constituation, anorexia, enteritis, flatulence; gastritis, jaundice, loose stools and oral monitiasis.

Hematologic and Lyggonatic: Anemia and leukopenia Nervous System: Headache (olilis media dosage), hyperkinesia, dizziness, agitation, nervousness and insomnia.

General: Fever, face edema, fatigue, fungal infection, malaise and pain.

insomnia.

General: Fever, face edema, fatique, tungal infection, malaise and pain.

Altergic: Rash and altergic reaction.

Respiratory: Cough increased, pharyogitis, pleural effusion and rhinitis.

Skin and Appendages: Eczema, tungal dermatitis, pruritus, sweating, urticaria and vesiculobullous rash.

Special Senses: Conjunctivitis.

Post-Markeling Expiritures

Adverse events reported with azithromycin during the post-marketing period in adult and/or pediatric palients for which a causal relationship may not be established include:

Altergic: Arthratija. edema, urticaria and angioedema.

Cardiovascular: Arthrytamias: including ventricular tachtycardia and hypotension. There have been rare reports of CTI proiongalion and torsades de pointes.

Gastrointestinai: Anotexia, coostigation, oxysepsia, liatulence and vomiting/diarrhea rarely resulting in chehydration, pseudomembrianous colitis, pancreabits, oral candidassis and rare reports of tongue discoloration General: Astheria, paresthesia, tallque, makisse and naphylosia (rarely tatal).

General: Astheria, paresthesia, fulloue, makise and naphylosia (rarely tatal).

General: Astheria presethesia, dulture, some of which have resulted in death.

Hematepo lettic: Thrombocytopenia;

Liver/Billiary: Abnormal liver function includion, hepatitis and cholestatic, jaundice, as well as rare cases of hepatic necrosis and hepatic faulture, some of which have resulted in death.

Hervous Systam: Connulsions, dizziness/vertigo, headache, sommolence, hyperactivity, nervousness, agitation and syncope.

agnation and synthyse. Psychiatric Repressive reaction and anxiety. Skin/Appendages: Printins, rarely serious skin reactions including crythema multiforme, Slevens Johnson Syndrome and toxic epidermal necrotysis.

Special Senses: Hearing disturbances including hearing loss, deafness and/or tinnitus and rare reports of taste perversion.

Issis perversion.

Laboratory Abnormalities:

Chincile Spiniticani abnormalities (irrespective of drug relationship) occurring during the clinical trials were challed.

Chincile Spiniticani abnormalities (irrespective of drug relationship) occurring during the clinical trials were challed to the control of the control o

Pediatric Patients:

One, Three and Five Day Regimens

Laboratory data collected from comparative clinical trials employing two 3-day regimens (30 mp/kg or 60 mg/kg) in divided doses over 3 days), or two 5-day regimens (30 mg/kg or 60 mg/kg) in divided doses over 3 days). Or two 5-day regimens (30 mg/kg as or 60 mg/kg) in divided doses over 3 days), or two 5-day regimens (30 mg/kg as a similar for regimens or altithorough and allocations of 15% Laboratory data for patients receiving 30 mg/kg as a single dose are single center intal. In that frial, an absolute neutrophi count between 500-1500 cells/mm² was observed in 10/64 patients receiving 30 mg/kg as a single dose, 9/62 galents receiving 30 mg/kg as a single dose, 9/62 galents receiving 30 mg/kg are not 30 mg/kg as a single dose, 9/62 galents received so mg/kg and 6/62 comparator patients, two patients and absolute neutrophil count <500 cells/mm². (See DOSAGE AND ADMINISTRATION.)

In multiple-dose situated links involving approximately 4/700 pediatric patients, no patients discontinued therapy because of treatment-related laboratory abnormalities.

DOSAGE AND ADMINISTRATION
TO A STREET AND ADMINISTRATION OF THE STREET AND CLINICAL PHARMACOLOGY.)

Infection*	Recommended Dose/Duration of Therapy
Community-acquired pneumonia (mild severity) Pharyngitis/tonsillitis (second line therapy) Skin/skin structure (uncomplicated)	500 mg as a single dose on Day 1, followed by 250 mg once daily on Days 2 through 5.
Acute bacterial exacerbations of chronic obstructive pulmonary disease (mild to moderate)	500 mg OD x 3 days OR 500 mg as a single dose on Day 1, followed by 250 mg once daily on Days 2 through 5.
Acute bacterial sinusitis	500 mg Q0 x 3 days
Genital ulcer disease (chancroid)	One single 1 gram dose
Non-gonoccocal urethritis and cervicitis	One single 1 gram dose
Gonococcal urethritis and cervicitis	One single 2 gram dose

Azithromycin oral suspension can be taken with or without food.

Authoritych or at suspension can be taken with or whologt 1000.

Renal lastificiency:

No dosage adjustment is recommended for subjects with renal impairment (GFR \$60 mL/min). The mean AUG\_1224 rate similar in subjects with GFR 10-80 mL/min compared to subjects with normal renal function, whereas it increased 35% in subjects with GFR <10 mL/min compared to subjects with normal renal function. Described should be subjects with normal renal function. Caution should be exercised when azimtomycin is administered to subjects with severe renal impairment. (See CLINICAL PHARMACOLOGY, Special Populations, Renal Insafficiency.)

Hepatic Instritutioners:

The pharmacobinetics of azithromyoin in subjects with hepatic impairment have not been established. No
The pharmacobinetic of azithromyoin in subjects with hepatic impairment have not been established. No
PHARMACOLOGY, Special Populations, Hepatic impairment have not been established. No
dosage adjustment is recommended based on age or gender. (See CLINICAL PHARMACOLOGY, Special
POpulations.)

Pediatric Patients: Azithromusic

reviews: Faterius:
Altithorogical asspension can be taken with or without tood.
Acute Dittis Media: The recommended dose of azithorogical or oral suspension for the treatment of pediatric patients with acute dittis media is 30 mg/kg given as a single dose or 10 mg/kg once daily for 3 days or 10 mg/kg as a single dose on the first day followed by 5 mg/kg/day on Days 2 through 5. (See chart below.)

Actual Bacterial Situstifis: The recommended dose of azithromycin for oral suspension for the treatment of pediatric patients with a quite bacterial sinusities is 10 mg/kg once daily for 3 days. (See chart below.) Community-Acquired Presuments: The recommended dose of asthromycin for oral suspension for the treatment of pediatric gatients with community-acquired pneumonia is 10 mg/kg as a single dose on the first day followed by 5 mg/kg on Days 2 through 5. (See Chart below.)

PEDIATRIC DOSAGE GUIDELINES FOR OTITIS MEDIA, ACUTE BACTERIAL SINUSITIS AND COMMUNITY-

OTITIS MEDIA AND COMMUNITY-ACQUIRED PNEUMONIA: 5-Day Regimen* Dosing Celculated on 18 mg/kg/day Day 1 and 5 mg/kg/day Days 2 to 5.							
We	ight		p/5 mL		g/5 mL	Total mil per	Total mg per
Kg	Lbs.	Day 1	Days 2-5	Day 1	Days 2-5	Treatment Course	Treatment Course
5	11	2.5 mL (1/2 tsp)	1.25 mL (1/4 tsp)			7.5 mL	150 mg
10	22	5 mL (1 tsp)	2.5 mL (1/2 tsp)			15 mL	300 mg
20	44			5 mL (1 tsp)	2.5 mL (1/2 tsp)	15 mL	600 mg
30	66			7.5 mL (11/2 tsp)	3.75 mL ( <sup>3</sup> / <sub>4</sub> tsp)	22.5 mL	900 mg
40	88			10 mL (2 tsp)	5 mL (1 tsp)	30 mL	1200 mg
50 and	110 and			12.5 mL (2 <sup>1</sup> / <sub>2</sub> isp)	6.25 mL (11/4 tsp)	37.5 mL	1500 mg

bove above Effectiveness of the 3-day or 1-day regimen in pediatric patients with community-acquired pne

	OHIS		BACTERIAL SINUSITI		]*
		Dosing Cal	culated on 10 mg/kg/	day	
Weight		100 mg/5 mL	200 mg/5 mL	Total mi. per	Total mg per
Kç	Lbs.	Day 1-3	Day 1-3	Treatment Course	Treatment Course
5	11	2.5 mL ( <sup>1</sup> / <sub>2</sub> tsp)		7.5 mL	150 mg
10	22	5 mL (1 tsp)		15 mL	300 mg
20	44		5 mL (1 tsp)	15 mL	600 mg
30	66		7.5 mL (1 ½ tsp)	22.5 mL	900 mg
40	88		10 mL (2 tsp)	30 mL	1200 mg
50 and above	110 and above		12.5 mL (2 ½ (sp)	37.5 mL	1500 mg

\*Effectiveness of the 5-day or 1-day regimen in pediatric patients with acute bacterial sinusitis has not beer

		DTITIS MEDIA: (1-0		<del></del>
		ng Calculated on 30 m		
Weight		200 mg/5 mL	Total mi. per	Total mg pe:
Kg	Lbs.	Day 1	Treatment Course	Treatment Course
5	11	3.75 ml. ( <sup>3</sup> / <sub>4</sub> lsp)	3.75 mL	150 mg
10	22	7.5 mL (1 <sup>s</sup> / <sub>2</sub> tsp)	7.5 mL	300 mg
20	44	15 mL (3 tsp)	15 mL	600 mg
30	66	22.5 mL (4 <sup>1</sup> / <sub>2</sub> tsp)	22.5 mL	900 mg
40	88	30 mL (6 tsp)	30 mL	1200 mg
50 and above	110 and above	37.5 mL (7 <sup>1</sup> / <sub>2</sub> tsp)	37.5 mL	1500 mg

The safety of re-dosing azithromycin in pediatric patients who vomit after receiving 30 mg/kg as a single dose has not been established. In clinical studies involving 487 patients with acute ofthis media given a single 30 mg/kg dose of azithromycin, eight patients who vomited within 30 minutes of dosing were re-dosed at the same total dose.

The same twist object.

Pharyagilis/fonstillis: The recommended dose of azithromycin for children with pharyagitis/ton mg/kg once daily for 5 days. (See chart below.)

PEDIATRIC DOSAGE GUIDELINES FOR PHARYNGITIS/TONSILLITIS
(Age 2 years and above, see PRECAUTIONS—Pediatric Use.)
Based on Body Weight

		esing Calculated on 12	mg/kg/day for 5 days.	
1	Veighl	200 mg/5 mL	Total mL per	Total mg per
Kg	Lbs.	Day 1-5	Treatment Course	Treatment Course
В	18	2.5 mL (1/2 tsp)	12.5 mL	500 mg
17	37	5 mL (1 1sp)	25 mL	1000 mg
25	55	7.5 mL (1 <sup>1</sup> / <sub>2</sub> tsp)	37.5 mL	1500 mg
33	73	10 mL (2 tsp)	50 mL	2000 mg
40	88	12.5 mL (2 <sup>1</sup> / <sub>2</sub> tsp)	62.5 mL	2500 mg

Constituting instructions for aziltromycin oral suspension, 300, 600, 900, 1200 mg bottles. The table below indicates the volume of water to be used for constitution:

Total volume after constitution (azithromycin content) 15 mL (300 mg) 15 mL (600 mg) 22.5 mL (900 mg) 30 mL (1200 mg) Azithromycin concentration after constitution 100 mg/5 mL 200 mg/5 mL 200 mg/5 mL 200 mg/5 mL Amount of Amount of water to be added 9 mL (300 mg) 9 mL (600 mg) 12 mL (900 mg) 15 mL (1200 mg)

Shake well before each use. Oversized bottle provides shake space. Keep tightly closed.

After mixing, store suspension between 5°C (41°F) and 25°C (77°F) and use within 10 days. Discard after full dosing is completed.

NOW SUPPLIED: Azithromycin for oral suspension, USP after constitution contains a flavored pink suspension. Azithromycin for oral suspension, USP is supplied to provide 100 mg/5 mL or 200 mg/5 mL suspension in bottles as 104mm/s.

# Azithromycin contents per bottle 300 mg 600 mg 900 mg 1200 mg

See DOSAGE AND ADMINISTRATION for constitution instructions with each bottle type. Storage: Store dry powder at 20"-25"C (68"-7.7"F) [See USP Controlled Room Temperature]. Store constituted suspension between 5°C (41"F) and 25°C (7"F) and discard when full dosting is completed.

# CLINICAL STUDIES (See INDICATIONS AND USAGE and Pediatric Use.)

Pediatric Patients from the perspective of evaluating pediatric clinical trials, Days 11-14 were considered on-therapy evaluations because of the extended half-life of azithromycin. Day 11-14 data are provided for clinical guidance, Day 24-32 evaluations were considered the primary lest of cure employin.

# Acute Otitis Media Safety and efficacy using azithromycin 30 mg/kg given over 5 days

Protocol 1

Fortocol 1 plan, controlled clinical study of acute office media performed in the United States, azithromycin (10 mg/kg on Days 1 followed by 5 mg/kg on Days 2-5) was compared to amount influedavidanate polassishm (4-1). For the SSS patients who were evaluated for clinical efficacy, but do with the Day 10 visit was 88% for patients with other particular protocol and the Day 10 visit, the clinical success rate was 75% for azithromych and 71% for the CO11 of the CO110 agent.

The clinical success rate was 75% for azithromych and 71% for the CO110 agent.

for the control agent. In the safety analysis of the above study, the incidence of treatment-related adverse events, primarily gastrolniestinal, in all patients treated was 9% with azilthromycin and 31% with the control agent. The most common side effects were distribused was a 20thromycin vs. 20% control, vomitting (2% azithromycin vs. 7% control), and addominal pain (2% azithromycin vs. 5% control), and addominal pain (2% azithromycin vs. 5% control).

azithromycin vs. 7% control), and adocument pain (2% azintromycin vs. 5% control).

Protocol 2
In a non-comparative clinical and microbiologic trial performed in the United States, where significant rates of beta-lactamase producing organisms (55%) were lound, 131 patients were evaluable for clinical efficacy. The combined clinical success rate (i.e., cure and improvement) at the Day 11 visit was 84% (iii) azintromycin: for the 122 patients who were evaluated at the Day 30 visit, the clinical success rate was 70% to azintromycin: for the 124 patients who were evaluated at the Day 30 visit, the clinical success rate was 70% to azintromycin: certain the comparation of the production of the

## Presumed Bacterlologic Eradication

S. pneumoniae H. influenzae M. catachalis S. pyogenes Overali	Day 11 Azithromycin 61/74 (82%) 43/54 (80%) 28/35 (80%) 11/11 (100%) 17//217 (82%)		Day 30 Azithromycin 40/56 (71%) 30/47 (64%) 19/26 (73%) 7/7 97/137 (73%)
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In the safety analysis of this study, the incidence of treatment-related adverse events, primarily gastrointestinal, in all patients treated was 9%. The most common side effect was diarrhea (4%).

gastrointestinal, in all patients treated was 9%, The most common side effect was disrribes (4%). Protocol 3
In another controlled comparative clinical and microbiologic study of obtis media performed in the United States, azithromycin was compared to amovicilitin/clavulanate polassium (4:1). This study villized two of the same investigators as Protocol 2 (above), and these two investigators enrolled 90% of the patients of beta-lactamase producing organisms (20%) were found. Minety-two (92) patients were evaluable tor clinical and microbiologic effictor. The combined clinical success rate (i.e., cure and improvement) of those patients with a baseline pathogen at the Day 11 visit was 80% for arithromycin vs. 100% for control; at the Day 30 visit, the clinical success rate (i.e., cure of the combined clinical success rate (i.e., of the control.) Microbiologic determinations were made at the pre-treatment visit. Microbiology was not reassessed at later visit. At the Day 11 and Day 30 visits, the clinical success she were obtained from the evaluable group:

## Procurant Racteriologic Fradication

.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Day	111	Day 30		
S. pneumoniae H. influenzae M. calarrhalis S. pyogenes Overall	Azithramycin 25/29 (86%) 9/11 (82%) 7/7 2/2 43/49 (88%)	Control 26/26 (100%) 9/9 5/5 5/5 45/45 (100%)	Azithromycin 22/28 (79%) 8/10 (80%) 5/5 2/2 37/45 (82%)	Control 18/22 (82%) 6/8 2/3 4/4 30/37 (81%)	

In the safety analysis of the above study, the incidence of treatment-related adverse events, primarily gastrointestinal, in all patients treated was 4% with aztimomycin and 31% with the control agent. The most common side effect was diarrhandose stools (5% aztimomycin x 25% control). Safety and efficacy asing azithromycin 30 mg/kg given over 3 days

Protocol 4
In a double-blind, controlled, randomized clinical study of acute othis media in pediatric patients from 6
months to 12 years of age, azithromycin (10 mg/kg per day for 3 days) was compared to
amoxicilliho/davulanate potassium (7:1) in divided doses q12h for 10 days. Each patients received active drug
and placebo matched for the comparator.
For the 366 patients who were evaluated for clinical efficacy at the Day 12 visil, the clinical success rate (i.e.,
cure plus improvement) was 38% for azithromycin and 88% for the control agent. For the 362 patients who
were evaluated at the Day 24-28 visil, the clinical success rate was 74% for azithromycin and 69% for the
control agent.

were evaluated at the Loy 24-cs val. the clinical success size was 24% for actinumyor and obey for the control algorit.

In the safety analysis of the above study, the incidence of treatment-related adverse events, primarily asstrointestinal, in all patients treated was 10.6% with azimonyoric and 20.0% with the control agent. The most common side effects were distributed abose stools (5.9% azimonyoric vs. 14.6% control), vomiting (2.1% azimonyoric vs. 1.1% control). Safety and efficacy using azilhremyoln 30 mg/kg given as a single dose.

Safety and efficacy using azilhremycin 30 mg/kg given as a single dose. 
Protocol 5
A double blind, controlled, randomized trial was performed at nine clinical centers. Pediatric patients from 6
norths to 12 years of age were randomized 1:1 to freatment with either aziltomonycin (given at 30 mg/kg as a single dose on Day 1) or amoxicillin/clavulantale potassium (7:1), divided q12h for 10 days. Each child received active drug, and placebo matches for the comparation.
Clinical response (Gure, Improvement, Failure) was evaluated at End of Therapy (Day 12-16) and Test of Cure (Day 26-20). Salety was evaluated throughout the trail for all treads obsjects, For the 321 subjects who were evaluated at End of Irreatment, the clinical success rate (cure plus improvement) was 87% for azithromycin and 88% for the comparation, for the 305 subjects who were evaluated at Test of Cure, the clinical success rate was 75% for both azithromycin and the comparation, the salety analysis, the incidence of treatment-related adverse events, primarity gastrointestinal, was 16,8% with azithromycin vs. 127% with the comparation. The most common side effects were diarrhea (6.4% with azithromycin vs. 127% with the comparator), common side effects were diarrhea comparation, was 16,8% with azithromycin vs. 1.2% with the comparator) and nauses (1.7% with azithromycin vs. 1.2% with the comparator).
Protocol 6
In a non-comparative clinical and microbiological trial. 248 patients from 6 months to 12 years of age with

Protection in a non-comparative clinical and microbiological trial. 248 patients from 6 months to 12 years of age with one name of the control of the contro

	Day 10	Day 24-28
S. pneumoniae	70/76 (92%)	67/76 (88%)
H. influenzae	30/42 (71%)	28/44 (64%)
M. catarrhalis	10/10 (100%)	 10/10 (100%)
Overall	110/128 (86%)	105/130 (81%)

Overall in the safety analysis of this study, the incidence of treatment-related adverse events, primarily gastrointestinal, in all the subjects treated was 12.1%. The most common side effects were vomiling (5.6%), and abdominal pair (1.6%), he most common side effects were vomiling (5.6%). Pharynghis/fore tillitis in the deather-bind contracted studies, conducted in the United States, azithromycin (12 mg/kg once a day for 5 days) was compared to pericilitin V (250 mg three times a day for 10 days) in the treatment of harynghis due to documented Group 4 h-themotyfic streptococci (6.68kg) or S. grogenes). Althomotymic was disincially and nicrobiologically statistically superior to pericilitin at Day 14 and Day 30 with the following clinical success, (it.e. cure and improvement) and bacteriologic efficacy rates (for the combined evaluable patient with documented GABHS):

Three U.S. Streptococcal Pharyngitis Studies
Azithromycin vs. Penicillin V
EFFICACY RESULTS
Day 14 Day 30

| Bacteriologic Eradication: Azithromycin | 323/40 (95%) | Penicilia V | Clinical Success (Cure plus improvement): | Azithromycin | 336/43 (98%) | Penicilia V | 284/338 (84%)

Approximately 1% of azithromycin-susceptible S. pyopenes isolates were resistant to azithromycin fo Approximately 1% to distinctive adverse events, primarily pastrointestinal, in all patients treated was 16% on patients of the primarily pastrointestinal, in all patients treated was 16% on a aztillomynch and 15% on penicillin. The most common side effects were dismeas/bose stools (5% aztillomynch ws. 1% penicillin), working (6% aztillomynch ws. 4% penicillin), and abdominal pain (3% adtillomynch ws. 1% penicillin). Active penicillini and abdominal pain (3% adtillomynch ws. 1% penicillini). Active penicillini and abdominal pain (3% adtill patients). Active Batterial Euscantalisms of Chronic Obstructive Pulmonary Oiscase in a randomated, double-bland controlled clinical trial or acute searchation of chronic bronchilis (AEC8), sathronneria (500 mg twice daily for 10

days). The primary endpoint of this trial was the clinical cure rate at Day 21-24. For the 304 patients analyzed in the modified intent to treat analysis at the Day 21-24 visit, the clinical cure rate for 3 days of azithromycin was 85% (12647) compared to 82% (1291/51) for 10 days of dariftmomycin. The following outcomes were the clinical cure rates at the Day 21-24 visit for the bacteriologically evaluable carrieds by exhibitons:

	Pathogen	•	Ċ	Azithromycin (3 Days)	Clarithromycin (10 Days)
S. pneumoniae H. influenzae			٠	29/32 (91%) 12/14 (86%)	21/27 (78%)
M. çatarrhalis				11/12 (92%)	14/16 (88%) 12/15 (80%)

M. catarnhalis 11/12 (92%) 12/15 (80%)
In the safety analysis of this study, the incidence of treatment-rebated adverse events, primarily opstrointestinal, were comparable between treatment arms (25% with azithromycin and 25% with carthromycin). The most common side effects were distribes, nauses and abdominal pain with comparable incidence rates for each symptom of 54% between the two treatment arms. (See ADVERS REACTIONS, ) Acute Bacterial Sinusilis in a randomized, double blind, double-dummy controlled clinical strial of acute-bacterial sinusilis azithromycin (500 mp. once daily for 3 days) was compared with amostellin/calvulantet (500/125 mp. lid for 10 days). Direction of the strial of acute-bacterial sinusilis azithromycin (500 mp. once daily for 3 days) was compared with amostellin/calvulantet (500/125 mp. lid for 10 days). Direction of the strial vote of the strial of the stria

of amodicilis/disvolanets.

In the safety analysis of this study, the overall incidence of treatment-related adverse events, primarily pastrointestinal, was lower in the azith/ornycin treatment arm (31%) than in the amodicilis/disvolanet arm (51%). The most common side electes were dearrheed (17% in the azith/ornycin arm vs. 32% in the amodicilis/disvolanete arm, and nausea (7% in the azith/ornycin arm vs. 12% in the amodicilis/disvolanete arm), (see ADVERSE REACTIONS).

In an open label, noncomparative sludy requiring baseline transantral sinus punctures the following outcomes vere the chinical success rates at the Day 7 and Day 25 wists for the modified intent to treat patients administered 500 mg of azithromycin once daily for 3 days with the following pathogens:

Pathogen			ycin or 3 Days)	
			Day 7	Day 28
o pneumoniae			23/26 (88%)	21/25 (84%
i, influenzae		٠.	28/32 (67%)	24/32 (75%

The overall incidence of treatment-related adverse events in the noncomparative study was 21% in modified intent to I read patients related with arthromych at 300 mg once daily for 3 days with the most common side effects being diarrhea (9%), abdominal pain (4%) and nausea (3%), (See AUPERS ERACTIONS).

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ANIMAL TOXICOLOGY
Phospholiolosis (intracellular phospholipid accumulation) has been observed in some tissues of mice, rats, and dogs given multiple doses of azilhromycin. It has been demonstrated in numerous organ systems (e.g., eye, dorsal root ganglia; liver, gallbadder, kidney, spleen, and pancreas) in dogs treated with azilhromycin at doses which, respected to the proposed of the proposed o

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