הודעה על החמרה (מידע בטיחות)

תאריך: 15.5.2012

שם תכשיר באנגלית: Invanz

מספר רישום: 127 09 30612 00

שם בעל הרישום: Merck, Sharp & Dohme (Israel-1996) Company Ltd.

השינויים בעלון <mark>מסומנים על רקע צהוב</mark>

בעלון לרופא

פרטים על השינוי/ים המבוקש/ים		
טקסט חדש	טקסט נוכחי	פרק בעלון
During clinical investigations in adult patients treated with INVANZ (1 g once a day), seizures, irrespective of drug relationship, occurred in 0.5% of patients during study therapy plus 14-day follow-up period [see Adverse Reactions (6.1)]. These experiences have occurred most commonly in patients with CNS disorders (e.g., brain lesions or history of seizures) and/or compromised renal function. Close adherence to the recommended dosage regimen is urged, especially in patients with known factors that predispose to convulsive activity. Anticonvulsant therapy should be continued in patients with known seizure disorders. If focal tremors, myoclonus, or seizures occur, patients should be evaluated neurologically, placed on anticonvulsant therapy if not already instituted, and the dosage of INVANZ re-examined to determine whether it should be decreased or discontinued.	Case reports in the literature have shown that co-administration of carbapenems, including ertapenem, to patients receiving valproic acid or divalproex sodium results in a reduction in valproic acid concentrations. The valproic acid concentrations may drop below the therapeutic range as a result of this interaction, therefore increasing the risk of breakthrough seizures. Increasing the dose of valproic acid or divalproex sodium may not be sufficient to overcome this interaction. The concomitant use of ertapenem and valproic acid/divalproex sodium is generally not recommended. Antibacterials other than carbapenems should be considered to treat infections in patients whose seizures are well controlled on valproic acid or divalproex sodium. If administration of INVANZ is necessary, supplemental anti-convulsant therapy should be considered.	WARNINGS- Seizure Potential
acid regurgitation, oral candidiasis, leg pain, anxiety, coordination abnormal, muscular weakness, gait disturbance		ADVERSE REACTIONS