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WARFARIN SODREM - warfarin sodium tablet
Physicians Tutal Care, Inc.
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Wastarin codium may increase the activated partial thromboplassin time (aPTT) two, even in the absence of heparin. A severe relevation (> 50 seconds) in aPTT with an INR in the desired range has been identified an analofaction of its revened risks of nonconcative beneathase.

Other Anticogulates:
Consult the labeling of other anticogulates for instructions on conversion to warfarineedism.

BOSAGE FORMS AND STRENGTHS
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nors not remnant air rate in cereming. Dings, dietary changes, and other factors affect INR levels achieved with warfarin sodium therape. Performance frequent INR measuring when starting or stopping other drugs, including bonaicals, or when changing docuges of other drugs [see Ding Interactions (7)].

Instruct patients about prevention measures to minimize risk of bleeding and to report signs and symptoms of bleeding [see Patient Counseling Information (17)]. 5.2 Tissue Necresia

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7.2 Drugs that Increase Bleeding Risk Examples of drugs known to increase the risk of bleeding are presented in Table 3. Recasse bleeding risk is increased when these drugs are used concontantly with warfarin, closely monitor patients receiving any each drug with warfarin.

	Table 3Drugs that Can Increase the Risk of Bleeding
Drug Class	Specific Drugs
ticoagulants	ogazuban, dabigazan, bivaliradin, desinadin, beparin, lepinadin
tiplatelet Agests	espiris, cilostatol, ciopidogre1, dipyridamole, prasugrei, siclopidine
ocuccidal Anti-	celecoxib, diciofesac, diffusical, fesoprafes, ibupcofes, indomethacis, ketoprofes, ketorulac, mefesamic acid, supr
lammatory Agents	

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Of the total number of gutients receiving wardarin sodium in controlled clinical trials for which data were available for analysis, title? patients (24.0%) were 65 years and older, while tile patients (2.0%) were 75 years and older. No overall differences in effectioness or safety were observed between these nations and control safety, but entered evolution of come older individuals or now he residences. were person that younger passes, and gravar centrality of storm sold for individuals, camer be relabelled.

The fillenth of juster or deling passes are shalling grown trappered DR requires the anticongular and passes of the control of the control

8.7 Repute Impairment
Hepatic Impairment can potentiate the response to warfarin through impaired synthesis of closing
factors and decreased methodisms of warfarin. Use castion when using warfarin codium in these
patients.

98. OVEROSAGE

10.1 Signs and Symptoms

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Hereding (e.g., appearance of the principles of the state of the support of the state of the support of the state of the st

The meatment of excessive anticangulation is based on the level of the INR, the presence or absence of bleeding, and clinical icinomentees. Reserval of warfaris sodium anticognistion may be obtained by discontinuing warfarin sodium therepy and, if one-ensury, by administrations of our of parenteral vibral



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uiled clinical trials involving 2711 patients with non-theustatic risk of symmic thromboetholismincluding stroke (see Table % to 86% in all except one trial (CAFA: 65%), which was e-results from two of these trials. The incidence of major 65% to 2.7% (see Table 4).

committee Enhage and Ammissionen (2.23).

18.2 Mechanical and Rieparcheie Heart Values
Ina prospector, makenized, open-bibel, positive-contribed undy in 254 gatieun with mechanical
psouthert heart valves, the threatherthelie-free inserval was found to be significantly goosen in
painten mound with variation above companyed with dispotant-polynimous and patients (p-0.005) and

Table SProspection,	Randomized, Ope Patients with M	n-Label, Pacitice-Councilled: lechanical Proothetic Heart V.	Clinical Study of Warfasia is slives
		Patients Treated With	
Event	Warfaria	Dipyridamale/Aspirin	Protectlylline/Aspirin
Thromboembolium	1.2/100 py	LE-1000 py	7.6r300 py

	Table 6Prospective, Open-Label Clinical Study of Martarin in Patients with Mechanic Prosched: Heart Values					
1	Event	Moderate	Marfaria Therapy	INR 2.65 High Intensity	Warfarin Therapy	INR S
ī	bromboembolism	47000 pv		3.7/1000 av		

including an analysis of st Tai	sscular death, are prov ble 7WARIS – Endoc		rate Events	
Event	Wartaria (N=607)	Placeba (N=607)	RR (95% CI)	% Risk Reduction (e-value)
Total Patient Years of Follow-up	2018	1944		
Total Mortality	94 (4.7/300 py)	123 (6.3/100 pg)	8.76 (0.60, 0.97)	24 (p~0.030
Vascular Death	82 (4.1/000 py)	105 (5.4100 py)	0.78 (0.60, 1.02)	
Recurrent MI	82 (4.1/000 py)	124 (6.4100 ps)	0.66 (0.51, 0.85)	34 (p~0.000

Major Bleeding* Minor Bleeding*	9	22	28	3.35 ^b (ND)	
Minor Bleeding ⁴				4.00° (ND)	ND ND
	29	103	133	3.21 th (ND) 2.50 th (ND)	ND ND
Composite Endpoints*	241	203	181	0.91 (0.60-0.95)* 0.71 (0.60-0.93)*	0.001
Reinfaccion	117	90	69	0.56(0.41-0.78)	0.03
Thromboembolic Smoke	20	17	17	0.52 (0.29-0.98)* 0.52 (0.29-0.97)*	0.03

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Warfarin Sodium Tables: USP, 7.5 mg R_s only





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