Low Vitamin D May Elevate Hypertension Risk

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NEW YORK — People with low serum levels of vitamin D have an increased risk of developing hypertension, based on the results from two prospective studies, one of which included more than 38,000

On the basis on both analyses, "we conclude that serum levels of 25-hydroxy vitamin D may be an independent risk factor for incident hypertension," Dr. John P. Forman said at the annual meeting of the American Society for Hypertension.

The findings also suggested that people could probably substantially reduce their risk by boosting their sun exposure through an extra 30-60 minutes spent outdoors daily, said Dr. Forman, a nephrologist at Brigham and Women's Hospital in Boston.

"Diet contributes relatively little to vitamin D levels, compared with sun exposure," said Dr. Forman. A typical multivitamin contains 200-400 IU of vitamin D. In contrast, an extra 30-60 minutes of sun exposure to the face and arms can generate 4,000 IU of vitamin D, although this relationship varies based on latitude and time of year. (Endogenous vitamin D production is minimal for people in the northern latitudes of the United States during the late fall and winter.)

Vitamin D deficiency has been hypothesized to cause hypertension based on epidemiologic studies that have shown that the further a population lives from the equator the higher the prevalence of hypertension. And mechanistic links exist: Active vitamin D suppresses renin expression in the juxtaglomerular apparatus, and vitamin D also inhibits the growth of vascular smooth muscle cells.

Both analyses used data collected from men enrolled in the Health Professionals Follow-up Study, which began in 1986. The first analysis focused on serum vitamin D levels measured from single blood samples drawn from 621 men who were normotensive at the time of their blood draw. These men had served as controls for a prior study that examined the link between vitamin D levels and the incidence of certain cancers.

In a multivariate analysis that controlled for a variety of demographic and clinical

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factors, including age, race, family history of hypertension, smoking, and alcohol intake, men with vitamin D deficiency (defined as a serum level of less than 15 ng/mL) had a statistically significant, 5.6fold increased risk of develop-

ing hypertension over the subsequent 4 years, compared with men whose baseline vitamin D level was 15 ng/mL or greater. During 8 years of follow-up, men with a baseline deficiency had a threefold increased risk of developing hypertension, although this difference was not statistically significant.

This analysis was limited by the relatively small number of men involved and because the reference vitamin D level was based on a single blood draw. Therefore, Dr. Forman and his associates ran a second analysis, based on data collected on all 38,388 men in the overall study.

Blood samples were not available for all of these men, but other available information allowed the researchers to estimate the subjects' serum level of vitamin D at baseline, using factors such as residence location, body mass index, and reported level of physical activity. The estimated vitamin D levels at baseline were correlated with the incidence of hypertension during the follow-up period, with adjustment for demographic and clinical

The analysis showed that men in the lowest decile for vitamin D had a 2.3-fold increased risk of new-onset hypertension, compared with men in the highest

The relationship between vitamin D and hypertension was biphasic. The relative risk gradually rose from the decile with the highest vitamin D level to a risk that was about 60% increased for the eighth decile. The risk then increased sharply for the two deciles with the lowest vitamin D levels.

CHANTIX (varenicline) TABLETS

INDICATIONS AND USAGE
CHANTIX is indicated as an aid to smoking cessation treatment.

PRECAUTIONS

General Nausea was the most common adverse event associated with CHANTIX treatment. Nausea was generally described as mild or moderate and often transient; however, for some subjects, it was persistent over several months. The incidence of nausea was dose-dependent. Initial dose-tituation was beneficial in reducing the occurrence of nausea. Nausea was reported by approximately 30% of patients treated with CHANTIX 1 mg BID dart an initial week of dose tritation. In patients taking CHANTIX 10.5 mg BID, the incidence of nausea was 16% following initial titration. Approximately 3% of subjects treated with CHANTIX 1 mg BID in studies involving 12 weeks of treatment discontinued treatment prematurely because of nausea. For patients with intolerable nausea, dose reduction should be considered. Effect of smoking cessation. Physiological changes resulting from smoking cessation, with or without treatment with CHANTIX, may after the pharmacokinetics or pharmacodynamics of some drugs, for which dosage adjustment may be necessarly (examples include theophylline, warrian and insulin.) (examples include theophylline, warfarin and insulin).

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Drug Interactions Based on verneicline characteristics and clinical experience to date, CHANTIX has no clinically meaningful pharmacokinetic drug interactions (See Full Prescribing Information, CLINICAL PHARMACOLOGY, Drug-Drug Interactions).

pnammazouristic origi mieracionis (see ruin resconnig miormation, cunicia. PHARMACULORY, Drug-Drug interactionis). Carcinogenesis, Mutagenesis, Impairment of Fertility. Carcinogenesis, Lifetime accinogenicity studies were performed in CO-1 mice and Sprague-Dawley rats. There was no evidence of a carcinogenic effect in mice administered varenicine by oral gavage for 2 years at doses up to 20 mpkg/ldg/4 (Pf times the maximum recommended human daily exposure based on ALIC). Rats evere administered varenicine brown fait were increased at the mid dose (I tumor, 5 mpkg/ldg/4, 23 times the maximum recommended human daily exposure based on ALIC). And maximum dose (2 tumors, 5 mpkg/ldg/4, 25 times the maximum recommended human daily exposure based on ALIC). The clinical relevance of this finding to humans has not been established. There was no evidence of carcinogenicity in female rats.

Mutagenesis. Varenicline was not genotoxic, with or without metabolic activation, in the following assays, sme bacterial mutation assay; mammalian CHO/HGPRT assay; and tests for cytogenetic aberrations in vivo in rat bone marrow and in vitro in human lymphocytes. Impairment of fertility. There was no evidence of impairment of fertility in either male or female Sprague-Dawley rats administered varenicline succinate up to 15 mg/kdyd y(67 and 68 times, respectively, the maximum recommended human daily exposure based on AUC at 1 mg BID). However, a decrease in fertility was noted in the offspring of pregnant rats who were administered varenicline succinate at an oral dose of 15 mg/kdy (65 times the maximum recommended human daily exposure based on AUC at 1 mg BID). This decrease in fertility in the offspring of treated female rats was not evident at an oral dose of 3 mg/kg/day (9 times the maximum recommended human daily exposure based on AUC at 1 mg BID).

fertility in the offspring of freated female rats was not evident at an oral dose of 3 mg/kg/day (9 times the maximum recommended human daily exposure based on AUC at 1 mg BID).

Pregnancy Pregnancy Category C. Varenicline succinate was not teratogenic in rats and rabbits at oral doses up to 15 and 30 mg/kg/day, respectively (36 and 50-times the maximum recommended human daily exposure based on AUC at 1 mg BID, respectively. Morteratogenic effects Varenicine succinate has been shown to have an adverse effect on the febus in animal reproduction studies. Administration of varenicine succinate to pregnant rabbits resulted in reduced fetal weights at an oral dose of 30 mg/kg/day (36 miss the production studies. Administration of varenicine succinate to pregnant rabbits resulted in reduced fetal weights at an oral dose of 30 mg/kg/day (36 miss the maximum recommended daily human exposure based on AUC). In addition, in the offspring of pregnant rats breated with varenicines succinate there were decreases in fertility and increases in auditory startle response at an oral dose of 15 mg/kg/day (36 times the maximum recommended human daily exposure based on AUC at 1 mg BID). There are no adequate and well-controlled studies in pregnant women. CHAITIX should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Nursing mothers: Although it is not known whether this darig is excreted in human milk and because of the potential for serious adverse reactions in nursing infants from CHAITIX, adverse and the serious adverse reactions in nursing pups. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing pups. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from GHAITIX, adverse to constitue from the potential for serious adverse reactions in the potential for serious adverse reactions in the potential for serious adverse reactions in nursing pups. B

- ormation for Patients:

 Patients should be instructed to set a date to quit smoking and to initiate CHANTIX treatment one week before the quit date.

 Patients should be advised that CHANTIX should be taken after eating, and with a full glass of water.

 Patients should be instructed how to titrate CHANTIX, beginning at a dose of 0.5 mg/day, Prescribers should explain that one 0.5 mg tablet should be taken daily for the first three days, and that for the next four days, one 0.5 mg tablet should be taken in the evening.

 Patients should be advised that, after the first seven days, the dose should be increased to one 1 mg tablet in the morning and one
- reactions should be advised that, after the miss seven days, the trues about the mideased where the religion in the miniming and the fing fable in the evening.

 Patients should be encouraged to continue to attempt to quit if they have early lapses after quit day.

 Patients should be informed that nausea and insomnia are side effects of CHANTIX and are usually transient; however, patients should be advised that if they are persistently troubled by these symptoms, they should notify the prescribing physician so that a dose effects of the properties of the properties

be advised that if they are persistently troubled by these symptoms, they should notify the prescribing physician so that a dose reduction can be considered.

• Patients should also be provided with educational materials and necessary counseling to support an attempt at quitting smoking.

• Patients should be informed that some medications may require dose adjustment after quitting smoking.

• Patients intending to become pregnant or planning to breast-feed an infant should be advised of the risks of smoking and risks and benefits of smoking cessation with CHANTIX.

ADVERSE REACTIONS

During the premarketing development of CHANTIX, over 4500 individuals were exposed to CHANTIX, with over 450 treated for at least 24 weeks and approximately 100 for a year. Most study participants were treated for 12 weeks or less. In Phase 2 and 3 placeborontrolled studies, the treatment discontinuation rates for the most common adverse events in CHANTIX placebo in studies of three months' treatment. In this group, the discontinuation rates for the most common adverse events in CHANTIX testop adjects were as follows: nauses (allows, 1,2% or placebo), Adverse Events were categorized using the Medical Dictionary for Regulatory Activities (MedDRA, Version 7.1).

The most common adverse events associated with CHANTIX (5-5% and twice the rate seen in placebo-treated patients) were nausea.

most common adverse events associated with CHANTIX (>5% and twice the rate seen in placebo-treated patients) were nausea, o disturbance, constipation, flatulence, and vomiting. Smoking cessation, with or without treatment, is associated with nicotine

wuuruawat syrupums.

The most common adverse event associated with CHANTIX treatment is nausea. For patients treated to the maximum recommended dose of 1 mg BID following initial dosage titration, the incidence of nausea was 30% compared with 10% in patients taking a comparable placebo regimen. In patients taking CHANTIX 0.5 mg BID following initial titration, the incidence was 16% compared with 11% for placebo. Nausea was generally described as mild or moderate and often transient, however, for some subjects, it was persistent throughout the treatment period.

Table 3 shows the adverse wants for CHANTIX and placebo in the 10 mild find of the compared to the com

Table 3 shows the adverse events for CHANTIX and placebo in the 12 week fixed dose studies with titration in the first week (Studies Cititated arm only), 4, and 5). MedDRA High Level Group Terms (HLGT) reported in ≥ 5% of patients in the CHANTIX 1 mg BID dose group, and more commonly than in the placebo group, are listed, along with subordinate Preterred Terms (PT) reported in ≥ 1% or CHANTIX patients (and at least 0.5% more frequent than placebo). Closely related Preterred Terms such as 'insonnia', 'Initia' insomnia', 'Hallde insomnia', 'Early morning awakening' were grouped, but individual patients reporting two or more grouped events are only counted once.

Table 3: Common Treatment Emergent AEs (%) in the Fixed-Dose, Placebo-Controlled Studies (≥1% in the 1 mg BID CHANTIX Group, and 1 mg BID CHANTIX at least 0.5% more than Placebo)

SYSTEM ORGAN CLASS High Level Group Term Preferred Term	CHANTIX 0.5 mg BID N=129	CHANTIX 1mg 1mg BID N=821	Placebo N=805
GASTROINTESTINAL			
GI Signs and Symptoms			
Nausea	16	30	10
Abdominal Pain*	5	7	5
Flatulence	9	6	3
Dyspepsia	5	5	3
Vomiting	1	5	2
Gl Motility/Defecation Conditions			
Constipation	5	8	3
Gastroesophageal reflux disease	1	1	0
Salivary Gland Conditions			
Dry mouth	4	6	4

(Table 3 continued)			
PSYCHIATRIC DISORDERS			
Sleep Disorders/Disturbances			
Insomnia**	19	18	13
Abnormal dreams		13	13 5 3
Sleep disorder	9 2 2	5	3
Nightmare	2	1 1	0
NERVOUS SYSTEM			
Headaches			
Headache	19	15	13
Neurological Disorders NEC			
Dysgeusia	8	5	4
Somnolence	3	3	2
Lethargy	2	1	0
GENERAL DISORDERS			
General Disorders NEC		_	
Fatigue/Malaise/Asthenia	4	7	6
RESPIR/THORACIC/MEDIAST			
Respiratory Disorders NEC			
Rhinorrhea	0	1 1	0
Dyspnoea	2	1 1] !
Upper Respiratory Tract Disorder	/	5	4
SKIN/SUBCUTANEOUS TISSUE			
Epidermal and Dermal Conditions			_
Rash] 3	4
Pruritis METABOLISM & NUTRITION	1 0		
Appetite/General Nutrit. Disorders	1 4		
Increased appetite	4	3	4

* Includes PTs Abdominal (pain, pain upper, pain lower, discomfort, tendemess, distension) and Stomach discomfort
** Includes PTs Insomnia/Initial insomnia/Middle insomnia/Early morning awakening

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The overall pattern, and the frequency of adverse events during the longer-term trials was very similar to that described in Table 3, though several of the most common events were reported by a greater proportion of patients. Nausea, for instance, was reported in 40% of patients treated with CHANTIX in mg BID in a one-year study, compared to 8% of placebo-treated patients.
Following is a list of treatment emergent adverse events reported by patients treated with CHANTIX during all clinical trials. The listing does not include those events which were so general as to be uninformative, and those events reported only once which did not awa a substantial probability of being acutely lite-threatening. BLOOD AND LYMPHATIC SYSTEM DISORDERS. Infrequent. Anemia, Lymphadenopathy. Rare. Leukocytosis, Thrombocytopenia, Splenomegaly, CARDIAC DISORDERS. Infrequent. Anemia, Lymphadenopathy. Rare. Leukocytosis, Thrombocytopenia, Splenomegaly, CARDIAC DISORDERS. Infrequent. Anemia, Lymphadenopathy. Rare. Leukocytosis, Thrombocytopenia, Splenomegaly, CARDIAC DISORDERS. Infrequent. Angina pectory. After Deathers, Merier's disease, Cor pulmonale, Acute coronary syndrome. EAR AND LABYRINTH DISORDERS. Infrequent. Thronic, Cardiac flutter, Coronary artery disease, Cor pulmonale, Acute coronary syndrome. EAR AND LABYRINTH DISORDERS. Infrequent. Conjunctivits, Dry eye, Eye irritation, Vision blurred, Visual disturbance, Eye pain. Rare. Acquired night blindness, Blindness transient, Cataract subcapsular, Ocular vascular disorder, Photophobia, Vitreous floaters, GASTROINTESTINAL DISORDERS. Frequent Diarrhea, Gingvitis, Infrequent: Cyslipacity, Cardia vascular disorder, Photophobia, Vitreous floaters, GASTROINTESTINAL DISORDERS Frequent Diarrhea, Gastric ulcer, Intestinal obstruction, Parceatitis, Event Increased, Infrequent: Hypersensitivity, Rar

DRUG ABUSE AND DEPENDENCE
Controlled Substance Class Varenicline is not a controlled substance. Humans: Fewer than 1 out of 1000 patients reported euphoria in clinical trials with CHANTIX. At higher doses (greater than 2 mg), CHANTIX produced more frequent reports of gastrointestinal disturbances such as nausea and vorniting. There is no evidence of doses-escalation to maintain therapeutic effects in clinical studies, which suggests that tolerance does not develop. Abrupt discontinuation of CHANTIX was associated with an increase in irritability and sleep disturbances in up to 3% of patients. This suggests that, in some patients, varencline may produce mild physical dependence which is not associated with addiction. In a human laboratory abuse liability subuy, a single oral dose of 1 mg varencline ind not produce which is not associated with addiction. In a human laboratory abuse liability subuy, a single oral dose of 1 mg varencline ind not produce any significant positive or negative subjective responses in morease in negative adverse effects, especially nausea. A single oral dose of 3 mg varencline uniformly produced unpleasant subjective responses in both smokers and non-smokers. Animals: Studies in roatina have shown that varencline produces behavioral responses similar to those produced by nicotine. In rats trained to discriminate nicotine from saline, varencline produced full generalization to the nicotine cue. In self-administration studies, the degree to which varencline from saline, varencline produced full generalization to the nicotine cue. In self-administration studies, the degree to which varencline continued to self-administer varencline to a degree comparable to that of nicotine, however in a more demanding task, rats self-administeralorine to a lesser extent than nicotine. Varencline pretreatment also reduced nicotine self-administration.

OVENUISME
In case of overdose, standard supportive measures should be instituted as required. Varenicline has been shown to be dialyzed in patients with end stage renal disease (see Full Prescribing Information, CLINICAL PHARMACOLOGY, Pharmacokinetics, Pharmacokinetics in Special Patient Populations), however, there is no experience in dialysis following overdose.

DOSAGE AND ADMINISTRATION

Usual Dosage for Adults Smoking cessation therapies are more likely to succeed for patients who are motivated to stop smoking and who are provided additional advice and support. Patients should be provided with appropriate educational materials and counseling to support the quit attempt. The patient should set a date to stop smoking, CHANTIX dosing should start one week before this date. CHANTIX should be taken after eating and with a full glass of water. The recommended dose of CHANTIX is 1 mg twice daily following a 1-week thatdom as follows:

Days 1-3:	0.5 mg once daily
Days 4-7:	0.5 mg twice daily
Days 8-End of treatment:	1 mg twice daily

Patients who cannot tolerate adverse effects of CHANTIX may have the dose lowered temporarily or permanently. Patients should be treated with CHANTIX for 12 weeks. For patients who have successfully stopped smoking at the end of 12 weeks, an additional course of 12 weeks treatment with CHANTIX is recommended to further increase the likelihood of long-term abstinence. Patients who do not succeed in stopping smoking during 12 weeks of initial therapy, or who relapse after treatment, should be encouraged to make another attempt once factors contributing to the failed attempt have been identified and addressed.

Special Populations

Patients with impaired renal function No dosage adjustment is necessary for patients with mild to moderate renal impairment. For patients with severe renal impairment, the recommended starting dose of CHANTIX is 0.5 mg once daily. Patients may then titrate as needed to a maximum dose of 0.5 mg once ally may be administered if tolerated well (See Full Prescribing Information, CLINICAL PHARMACOLOGY, Pharmacokinetics, Pharmacokinetics in Special Populations, Renal impairment.) Dosing in elderly patients and patients with impaired hepatic function. No dosage adjustment is necessary for patients with hepatic impairment. Because elderly patients and patients with impaired hepatic function. No dosage adjustment is necessary for patients with hepatic impairment. Because elderly patients and are role likely to have decreased renal function. Care should be taken in dose selection, and it may be useful to monitor renal function. Ges PBECAUTIONS, Geriatric Use). Use in children Safety and effectiveness of CHANTIX in pediatric patients have not been established; therefore, CHANTIX is not recommended for use in patients under 18 years of age.

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