Rising Anal Cancer Incidence Parallels HPV, AIDS

BY TIMOTHY F. KIRN Sacramento Bureau

SAN FRANCISCO — The HIV epidemic has brought to light evidence that men who have sex with men now have a rate of anal cancer as high as the incidence rate of cervical cancer in women prior to Pap smear.

The incidence rate in women also has risen, Dr. Joel Palefsky said at a meeting on HIV management sponsored by the

CHANTIX

(varenicline) tablets

PRECAUTIONS

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

University of California, San Francisco.

The incidence rate of cervical cancer prior to Pap smear screening was 40-50 cases per 100,000 females. The present rate of anal cancer in HIV-negative men who have sex with men is 35 per 100,000, and the rate for HIV-positive men is probably twice as high, said Dr. Palefsky, a professor of medicine and the director of the Anal Dysplasia Clinic at the UCSF Cancer Center.

Both anal dysplasia and cervical cancer are related to human papillomavirus (HPV) infection. The incidence rate in HIV-positive men appears to be rising for a few reasons. Highly active antiretroviral therapy (HAART) is prolonging the life of HIV-infected individuals long enough for cancer to develop. HIV infection itself accelerates the process, said Dr. Palefsky, who is part of group urging anal Pap smear screening for at-risk individuals. "If you are HIV positive, you have anal HPV infection." he said.

Studies he conducted in the pre-HAART

era found abnormal anal cytology in 80% of HIV-infected men with CD4 T-cell levels less than 200 cells/mcL, Dr. Palefsky said. Since then, studies have shown that HPV-infected persons develop intraepithelial neoplasia regularly and rapidly, and that HAART may not lower the incidence (Clin. Infect. Dis. 2002;35:1127-34).

Women too have high rates of anal HPV infection, and it is not just HIV-positive women, he said. In the Women's Interagency HIV Study, they found a high prevalence of HPV infection. In fact, anal HPV was more common than cervical infection in both 251 HIV-positive women (79% vs. 53%) and in 68 HIV-negative women (43% vs. 24%) (J. Infect. Dis. 2001;183:383-91).

Surveys he did in Planned Parenthood clinics and a cervical dysplasia clinic also suggest that anal HPV infection is as least as common as cervical HPV in women.

Evidence that anal cancer is becoming more common is less certain, Dr. Palefsky said. Data from the Surveillance, Epidemiology, and End Results survey suggest that squamous cell carcinoma of the anus has



Squamous cell carcinoma of the anus has increased perhaps 96% in men and 39% in women over the past 40 years.

DR. PALEFSKY

increased perhaps 96% in men and 39% in women over the past 40 years.

Much of anal HPV infection results from receptive anal intercourse, but not all. It has been found to be common in HIV-positive men who are injection drug users.

Data suggest that half of women in this country engage in anal intercourse at least once in their lifetime, and, "as we know from the cervix, it doesn't take too many exposures to get HPV," he noted.

Dr. Palefsky said he examines for anal dysplasia the same way one does for cervical dysplasia. He inserts a water-moistened Dacron swab into the anus and rubs the swab around the wall of the canal. The material obtained is graded according to the Bethesda System like a cervical sample.

A digital rectal exam, done after the swabbing, requires lubricant. "You will feel things with your finger that you'll not see either on a Pap smear or through the anoscope because a lot of the cancers are subcutaneous at that point," he said.

Patients with a positive Pap smear undergo examination with a cervical colposcope. A gauze pad is soaked in a vinegar solution (2:1 vinegar to water), then wrapped around a swab and inserted through an anoscope. The anoscope is then removed, and the gauze is left in place for 1 minute. Once the gauze is removed, the anoscope is reinserted and examination is done with the colposcope.

The same signs used for the cervix have been validated in the anal canal, he said.

Small lesions can be treated with trichloroacetic acid, and larger lesions can be treated with infrared coagulation, which is being shown to be successful, he added.

(Table 3 contin PSYCHIATRIC DISORDERS Sieep Disorders/Disorders/Disorders/Disorders/Disorders/Disorders/Disorder rs/Disturbances 18 13 Nightmare NERVOUS SYSTEM Headaches Headache Neurological Disorders NEC Dysgeusia Somnolence 19 15 13 Lethargy GENERAL DISORDERS General Disorders NEC Fatigue/Malaise/Astheni RESPIR/THORACIC/MEDIA espiratory Disorders Rhinorrhea 0 0 Upper Respiratory Tract Disorder SKIN/SUBCUTANEOUS TISSUE Epidermal and Dermal Condit Rash 3 2 Pruritis METABOLISM & NUTRITION

** Includes PTs Abdominal (pain, pain upper, pain lower, discomfort, tendeness, distension) and Stomach discomfort ** Includes PTs Insomnia/Initial insomnia/Initial insomnia/Early moming avakening he overall pattern, and the frequency of adverse events during the longer-term trials was very similar to that described in Table 3, hough several of the most common events were reported by a grateer propriotin of patients. Nausea, for instance, was reported in 0% of patients treated with CHANTIX 1 mg BD in a one-year study, compared to 8% of placebo-treated patients. Diolwing is a list of treatment-mergent adverse events reported by patients treated with CHANTIX during all clinical trials. The listing one not include those events fore apoint adverse events reported by patients treated with CHANTIX during all clinical trials. The listing one not include those events fore apoint adverse events uniformative and these specific exported for patients treated with compared to patients.

40% of patients treated with CHANILX I mg HU in a One-year study, compared to any to pracetou-treased patients. Following is a list of treatment-encegnet adverse serverits reported by patients treated with CHANTX during al clinical trials. The listing does not include those events already listed in the previous tables or elsewhere in labeling, those events for which a during cause was ender, those events which were so general as to be uninformative, and those events in proteot only once which did not have a substantial probability of being acutely life-threatening. BLOOD AND LYMPHATIC SYSTEM DISORDERS. Infrequent: Anemia, Lymphadenogathy, Rare, Leukoytosis, Thromboytopenia, Splenomegaju, CABNDAC DISORDERS. Infrequent: Anemia, Lymphadenogathy, Rare, Leukoytosis, Thromboytopenia, Splenomegaju, CABNDAC DISORDERS. Infrequent: Anemia, Lymphadenogathy, Rare, Leukoytosis, Thromboytopenia, Splenomegaju, CABNDAC DISORDERS. Infrequent: Throng and disorders. EVE DISORDERS. Infrequent: Conjunctive, Acute coronary synchrome. EAR AND LABYRNITH DISORDERS. Infrequent: Torolita, Puter States, Schemes events, Moinh were achesisted as the comparison of the second synchrone and disorders. EVE DISORDERS infrequent: Conjunctive, Schemes transmit, Verguent, Disorders, States Constructive, Through and disorders. EVE DISORDERS infrequent: Conjunctive, Not Useration, Espanja, Turking, Kare, Dug Popenagitis, Rare, Dug Popenagitis, Rare, Dug Popenagitis, Rare, Dug Popenagitis, Rare, Dug Popenasitivity, INVESTIGATIONS, Frequent Liver function test abrormal, Weight increased. Infrequent: Electroardiogram abnormal, Wiscale enzyme Increased, Universal, Miscale abrormal, Weight, Increased, Infrequent: Electroardiogram abnormal, Wiscale enzyme Increased, Universal, Publica Vester, IMMUNE SYSTEM DISORDERS, Infrequent: Electroardiogram abnormal, Wiscale enzyme Increased, Universal, Publica Vester, MINUNE SYSTEM DISORDERS, Infrequent: Electroardiogram abnormal, Miscale enzyme Increased, Internation, Disorderist, Indrequent, Rare, Mystis, NEFVOUS S

DRUG ABUSE AND DEPRODENCE Controlled Substance Class Varenicine is not a controlled substance. <u>Humans</u>: Fewer than 1 out of 1000 patients reported euphoria in clinical trials with CHAITN. At higher doses (greater than 2 mg), CHANTIX produced more frequent reports of gastrointestinal disturbances such as nausea and vomting. There is no evidence of dose-escalation to maintain therapeutic effects in clinical studies, which suggests that blearance does not develop. Abrupt discontinuation of CHAITIX was associated with an increase in initiability and sleep disturbances in up to 3% of patients. This suggests that, in some patients, varenicine may produce mild physical dependence which is not associated with addiction. In a human laboratory abuse lability study, a single eral dose of 1 my varenicine id not produce any significant positive or negative subjective responses in mons mokers. In mon-smokers, 1 my varenicine produced an oral dose of 3 my varenicine unformly produced unpleasant subjective responses in both smokers and non-smokers. <u>Animals</u>: Studies in rodents have shown that vernicine produced full generalization to the nicotine cue. In self-administration studies, the degree to which varenicine substitutes for incoline is dependent upon the requirement of that of incoline, however in a more demanding task, rads self-administer varenicine to a lesser extent than microtine. Varenicine produced minister incoline useft-administration. **OVERDOSAGE** OVERDOSAGE

VUTENUXANCE 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 PHARMACOLOCY, Pharmacokinetics, Pharmacokinetics in Special Patient Populations), however, there is no experience in dialysis following overdose.

Usual Dosage for Adults Smoking essation 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 titration as follows:

Days 4–7: 0.5 mg twice daily Days 8–End of treatment: 1 mg twice daily	Days 1-3:	0.5 mg once daily	
Days 8-End of treatment: 1 mg twice 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 limital 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 twice a day. For patients with End-stage renal disease undergoing hemodialysis, a maximum dose of 0.5 mg once daily may be administered if theirated well (See Tull Prescribing Information, CLINACL PHARIMACOLOGY, Pharmacokinetics, Pharmacokinetics in Special Populations, Renal impairment, Dosing in elderly patients and patients with severe renal impairment, and the severe of the severe of the severe renal disease endergoing hemodialysis, are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function. (See PECEL/UTIONS, Certairtu Use). Use in children Safety and effectiveness of CHANTIX is pediatric patients have not been established; therefore, CHANTIX is not recommended for use in patients under 18 years of age.

May 2006, Version LAB-0327-2.0

PRECAUTIONS General Nausea was the most common adverse event associated with CHANTX 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-titation was beneficial in reducing the occurrence of nausea. Nausea was reported by approximately 30% of patients treated with CHANTX. If ang BDI after an initial week of dose tritandin. In patients taking CHANTO. Os m BDI, the incidence of nausea was 15% following initial titration. Approximately 3% of subjects treated with CHANTX in gBID in studies involving 12 weeks of treatment discontinued treatment prematurely because of nausea. Are patients with inderable nausea, dose reduction should be considered. *Effect of smoking cessation*: Physiological changes resulting from smoking cessation, with or without treatment with CHANTX, may after the pharmacohicits or pharmacohymanics of some drugs, for which dosage adjustment may be necessary (examples include theophyline, warfarin and insulin). (examples include theophyline, waratin and insulin). Drug Interactions Based on varatin and insulin). Drug Interactions Based on varaticities the end of Mutagenesis. Varenicline was not genotoxic, with or without metabolic activation, in the following assays: Ames bacterial mutation assay; mammalian CHO/HGPRT assay; and tests for cytogenetic aberrations *in vivo* in rat bone marrow and *in vitro* in human lymphocytes.

Intellination Churnerria essey, and tess to d'oyderiete aderitations in vivo in ratio un terrativo and in vincon induitari organizopes. Impairment of fertility. There vas en evidence or impairment of fertility in either male or fernale Sprague-Dawley rats administreed varianciane succinate up to 15 mg/kg/day (62 and 36 times, respectively, the maximum recommended human daily exposure based on AUC at 1 mg BID), Howerer, a decrease in fertility was notevident in the offspring or program trats who were administreed varianciane succinate an oral dose of 15 mg/kg/day (68 times the maximum recommended human daily exposure based on AUC at 1 mg BID). This decrease in fertility in the offspring of treaded termale 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** Tergenary 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. Nontreatogenic effects Varenicine succinate was not teratogenic in rats and rabbits at oral doses up to 15 and 30 mg/kg/day, respectively. Montreatogenic effects Varenicine succinate has been shown to have an adverse effect on the fetus in animal reproduction studies. Administration of varencline succinate to pregnant rabbits resulted in reduced fetal veipths at an oral dose of 3 mg/kg/day (23 times the maximum recommended daily human exposure based on AUC). In addition, in the offspring of pregnant rab treated with varencline succinate has been advected to 15 mg/kg/day (36 times the maximum recommended daily human exposure based on AUC). In addition, in the offspring of pregnant rab treated with varencline succinate have were decreases in fulfily 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. CHANTIX should be used during pregnancy only if the potential benefit justifies the potential ins to the fusus. **Nursing mothers** Although its not known whether this during is excreted in human milk and because of the potential lor serious adverse reactions in nursing puse. Because many drugs are excreted in human milk and because of the potential lor serious adverse reactions in account the importance of the drug to the mother. **Labor and delivery** The potential effects O CHANTIX on labor and delivery are not known. **Pediatric Use** Safety and the reported dinical experience has not identifited mate and themas

ents (see DOSAGE AND AUMINISTIATION, opposed a second of initiate CHANTX treatment one week before the quit date. • Patients should be instructed how that CHANTX should be taken after eating, and with a full glass of water. • Patients should be instructed how to thate CHANTX, 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 worning and 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 1 mg tablet in the evening. • Determine the evening.

Parents structure downeed units, endowneed and the version of the version of the version.
Parents should be encouraged to continue to attempt to quit if they have early lapses after quit day.
Parents should be informed that nauses and insomma are side effects of CHANTX 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 reduction can be considered.

duction can be considered. distints should also be provided with educational materials and necessary counseling to support an attempt at quitting smoking, atients should be informed that some medications may require dose adjustment after quitting smoking. atients intending to become pregnant or planning to breast-feed an infant should be advised of the risks of smoking and risks and metits of smoking cessation with CHWITX.

ADVERSE REACTIONS

ADVERSE FEACTONS 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 placebo-controlled studies, the treatment discontinuation rate due to adverse events in patients dosed with 1 mg BID was 12% for CHANTIX compared to 10% for placebo in studies of three months' treatment. In this group, the discontinuation rates for the most common adverse events in CHANTIX treated patients were as follows: nausea (3% vs. 0.5% for placebo), headache (0.6% vs. 0.9% for placebo), insomnia (12.4% vs. 1.1% for placebo), and abnormal dreams (0.3% vs. 0.2% for placebo), Adverse Events were categorized using the Medical Dictionary for Regulatory Activities (MedDRA, Version 7.1).

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

withdrawal symptoms. 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. Ruusea 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 events for CHANTIX and placebo in the 12 week fixed dose studies with titration in the first week (Studies 2 (titrated arm only), 4, and 5). MedDRA High Level Group Terms (HLGT) reported in ≥ 5% of patients in the CHANTIX that may and more commonly than in the placebo group, are listed, along with subordinate Preferred Terms (PT) reported in ≥ 1% of ANTIX platents (and at least 0.0%) more frequent than placebo (Dosely related Preferred Terms such as "Insomnia", "Initial insomnia", "Middle insomnia", "Early morning awakening" were grouped, but individual patients reporting two or more grouped events

Table 3: Common Treatment Emergent AEs (%) in the Fixed-Dose, Placebo-Controlled Studies (\geq 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
GI Motility/Defecation Conditions			
Constipation	5	8	3
Gastroesophageal reflux disease	1	1	0
Salivary Gland Conditions			
Dry mouth	4	6	4

© 2007 Pfizer Inc.

VC276999G

All rights reserved.

Printed in USA/January 2007

Pfizer U.S. Pharma

Before prescribing, please consult Full Prescribing Information. Appetite/General Nutrit. Disorders Increased appetite Decreased appetite/Anorexia 4 3 2 2