HHS Aims to Spur Electronic Biosurveillance

BY MARY ELLEN SCHNEIDER

Senior Writer

WASHINGTON — Government officials and health information technology leaders plan to spend this year laying the groundwork for a system that would allow for the electronic transfer of ambulatory, emergency department, and laboratory data to public health agencies in less than a day.

Over time, officials would like to implement a real-time nationwide public health monitoring system. "The system we have is simply not adequate," Mike Leavitt, secretary of the Health and Human Services department, said at a meeting of the American Health Information

The United States faces not only the possibility of a bioterrorist attack but also the threat of pandemic, he said.

Mr. Leavitt said he would like to get a 'spotty net" of surveillance off the ground quickly by collecting a few key indicators from as many electronic data sources as possible. Getting just 2-4 basic data points from all available sources would be a quantum leap forward," he said.

Information from small and mediumsized primary care practices will be key to any electronic biosurveillance system, said Dr. David Kibbe, who represented the American Academy of Family Physicians at the meeting. The American Health Information Community is an advisory committee to the Health and Human Services department. "There is widespread agreement that information technology can substantially improve surveillance both for ongoing public health and for health emergencies," said Dr. Thomas R. Frieden, commissioner of the New York City Department of Health and Mental Hygiene, who presented information on current electronic surveillance programs at the meeting.

Biosurveillance activities are underway at the federal, state, and local levels, and in the private sector, Dr. Frieden said. For example, the Centers for Disease Control and Prevention operates the Public Health Information Network, which provides an architecture for public health information technology. Most recently, the agency established the BioSense program which is

Already, 50 New **York City** hospitals report data on a daily basis to the city health department on ED and outpatient visits, pharmacy

otherwise.

aimed at supporting the connection of clinical care to public health and supporting "situational awareness" at the national level. A number of state and local health departments have begun electronic reporting either

purchases, etc. from clinical laboratories or clinical information systems. In New York City, the health department uses electronic reporting data on a daily basis. The system, which has been operating for more than 5 years, collects information from ambulance dispatches, emergency department visits, pharmacy purchases, outpatient visits, and other sources. The system also collects free text, which allows officials to evaluate information they might not have thought about

Currently, 50 hospitals—representing about 90% of emergency department visits in the city—report daily.

The electronic reporting system has proved helpful in the early detection of pockets of influenza. The electronic syndromic system consistently picks up influenza activity 2-3 weeks before any other system.

New York City is not alone. North Carolina has a statewide, hospital-based clinical data monitoring system. It allows for monitoring of real-time inpatient, outpatient, and emergency department data.

But there are major needs that must be addressed to reach the goal of a nationwide system, Dr. John Loonsk of the federal Office of the National Coordinator for Health Information Technology said at the meeting.

For example, data need to be standardized so they can be compared across reporting organizations, privacy and confidentiality must be ensured, and improvements need to be made in the current patchwork of state and local health information technology capability, he said.

One area that potentially could be implemented rapidly is the electronic reporting of lab results, Dr. Loonsk said.

Brief Summary of Prescribing Information (Nos. 1541, 1543, 1544, 3046, 7309, 7311) n3-5366-R24-Brf. Rev. July, 2004

PREVACID® (lansoprazole) Delayed-Release Capsules

PREVACID® (lansoprazole) For Delayed-Release Oral Suspension

PREVACID® SoluTabTM (lansoprazole) Delayed-Release Orally

VACID Delayed-Release Capsules, PREVACID SoluTab Delayed-Release Orally ntegrating Tablets and PREVACID For Delayed-Release Oral Suspension are indicated

Triple Therapy: PREVACID/amoxicillin/clarithromycin
Dual Therapy: PREVACID/amoxicillin/clarithromycin or in whom resistance to
clarithromycin is known or suspected.
Maintenance of Healed Duodenal Ulcers
Controlled Studies do not extend beyond 12 months.

Controlled studies do not extend beyond 12 months.
Short-Term Treatment (up to 8 weeks) of Active Benign Gastric Ulcer
Healing of NSAID-Associated Gastric Ulcer
In patients who continue NSAID use. Controlled studies did not extend beyond 8 weeks.
Risk Reduction of NSAID-Associated Gastric Ulcer
In patients with a history of a documented gastric ulcer who require the use of an NSAID.
Controlled studies did not extend beyond 12 weeks.
Gastroesophageal Reflux Disease (GERD)
Short-Term Treatment of Symptomatic GERD)
Short-Term Treatment (up to 8 weeks) of Erosive Esophagitis
For patients who do not heal with PREVACID for 8 weeks (5-10%), it may be helpful to give an additional 8 weeks of terrosive Esophagitis an additional 8 weeks course of PREVACID may be considered.
Maintenance of Healing of Erosive Esophagitis

adultion of the control of the contr

Pathological Hypersecretory Conditions Including Zollinger-Ellison Syndrome
CONTRAINDICATIONS
PREVACID is contraindicated in patients with known hypersensitivity to any component of
the formulation of PREVACID.
Amoxicillin is contraindicated in patients with a known hypersensitivity to any penicillin.
Clarithromycin eyrthormycin, and any of the macrolide antibiotics.
Concomitant administration of clarithromycin with cisapride, pimozide, astemizole, or
terfenadine is contraindicated. There have been post-marketing reports of drug interactions
when clarithromycin and/or erythromycin are co-administered with cisapride, pimozide,
astemizole, or terfenadine resulting in cardiac arritythmisa (OT prolongation, ventricular
tachycardia, ventricular fibrillation, and torsades de pointes) most likely due to inhibition of
metabolism of these drugs by erythromycin and clarithromycin. Fatalities have been
reported.

Please refer to full prescribing information for amoxicillin and clarithromycin before prescribing.)

(Please refer to full prescribing information for amoxicillin and clarithromycin before prescribing.)

WARNINGS
CLARITHROMYCIN SHOULD NOT BE USED IN PREGNANT WOMEN EXCEPT IN CLINICAL CIRCUMSTANCES WHERE NO ALTERNATIVE THERAPY IS APPROPRIATE. IF PREGNANCY OCCURS WHILE TAKING CLARITHROMYCIN, THE PATIENT SHOULD BE APPRISED OF THE POTENTIAL HAZARD TO THE FETUS. (SEE WARNINGS IN PRESCRIBING INFORMATION FOR CLARITHROMYCIN.)

PSEUdomembranous collist has been reported with nearly all antibacterial agents, including clarithromycin and amoxicillin, and may range in severity from mild to life threatening. Therefore, it is important to consider this diagnosis in patients who present with diarnhea subsequent to the administration of antibacterial agents. Treatment with antibacterial agents alters the normal flora of the colon and may permit overgrowth of clostridia. Studies indicate that a toxin produced by Clostridium difficile is a primary cause of "antibiotic-associated collis".

After the diagnosis of pseudomembranous collis has been established, therapeutic measures should be initiated. Mild cases of pseudomembranous collis usually respond to discontinuation of the drug alone. In moderate to severe cases, consideration should be given to management with fluids and electrolytes, protein supplementation, and treatment with an antibacterial drug clinically effective against Clostridium difficile collis. Serious and occasionally fatal hypersensitivity and/or a history of sensitivity moditions when treated with a cephalosporin. Before initiating therapy with any penicillin, careful inquiry should be made concerning previous hypersensitivity reactions have been reported in patients on penicillim hypersensitivity and/or a history of sensitivity moditions when treated with a cephalosporin. Before initiating therapy with any penicillinic, careful inquiry should be made concerning previous hypersensitivity reactions have been reported in made concerning previous hypersensitivity reactions have been reported with a near of conce

uvereralSymptomatic response to therapy with lansoprazole does not preclude the presence of gastric malignancy.

gastric manigrancy.

Information for Patients
PREVACID is available as a capsule, orally disintegrating tablet and oral suspension, and is available in 15 mg and 30 mg strengths. Directions for use specific to the route and available in 15 mg and 30 mg strengths. Directions for use specific to the route and available methods of administration for each of these dosage forms is presented below. PREVACID should be taken before eating. PREVACID products SHOULD NOT BE CRUSHED OR CURLING.

Phenylketonurics: Contains Phenylalanine 2.5 mg per 15 mg Tablet and 5.1 mg per 30 mg Tablet.

30 mg Tablet.

Administration Options

1. PREVAIDID Delayed-Release Capsules

1. PREVAIDID Delayed-Release Capsules

PREVAIDID Delayed-Release Capsules should be swallowed whole.

Alternatively, for patients who have difficulty swallowing capsules, PREVACID Delayed-Release Capsules can be opened and administered as follows:

• Open capsule.

• Open capsule.

Sprinkle intact granules on one tablespoon of either applesauce, ENSURE® pudding, cottage cheese, yogurd or strained pears.

• Swallow immediately.

PREVACID Delayed-Release Capsules may also be emptied into a small volume of either apple juice, orange juice or tomato juice and administered as follows:

- Open capsule:

en capsule.

Inkle intact granules into a small volume of either apple juice, orange juice or tomato e (60 mL – approximately 2 ounces).

Specifically a small volume of either apple juice, orange juice or tomato e (60 mL – approximately 2 ounces).

Swallow immediately.
 Fo ensure complete delivery of the dose, the glass should be rinsed with two or more
volumes of juice and the contents swallowed immediately.
 USE IN OTHER FOODS AND LIQUIDS HAS NOT BEEN STUDIED CLINICALLY AND IS
THEREFORE NOT RECOMMENDED.

HEHEHORE NOT RECOMMENDED.

2. PREVAICID Soluriab Delayed-Release Orally Disintegrating Tablets

PREVAICID Soluriab should not be chewed. Place the tablet on the tongue and allow it to disintegrate, with or without water, until the particles can be swallowed. The tablet typically disintegrates in less than 1 minute.

Alternatively, for children or other patients who have difficulty swallowing tablets, PREVAICID Soluriab can be delivered in two different ways.

PREVACIO Solu Tab can be delivered in two different ways.

PREVACIO Solu Tab — Oral Syringe, PREVACIO Solu Tab can be administered as follows:

Palea a 15 mg lablet in oral syringe, PREVACIO Solu Tab can be administered as follows:

Place a 15 mg lablet in oral syringe and draw up approximately 4 mL of water, or place a 30 mg tablet in oral syringe and draw up approximately 10 mL of water.

30 mg tablet in oral syringe and draw up approximately 10 mL of water.

51 hate gently to allow for a quick dispersal.

After the tablet has dispersed, administer the contents within 15 minutes.

Felfill the syringe with approximately 2 mL (5 mL for the 30 mg tablet) of water, shake gently, and administer any remaining contents.

VACID SoluTab – Nasogastric Tube Administration (≥ 8 French) administration via a nasogastric tube, PREVACID SoluTab can be administered as

follows:

Place a 15 mg tablet in a syringe and draw up 4 mL of water, or place a 30 mg tablet in a syringe and draw up 10 mL of water.

Stake gently to allow for a quick dispersal.

After the tablet has dispersed, inject through the nasogastric tube into the stomach within 15 minutes.

Refill the syringe with approximately 5 mL of water, shake gently, and flush the nasogastric tube.

3. PREVACID for Delayed-Release Oral Suspension
PREVACID for Delayed-Release Oral Suspension should be administered as follows:

perspaces.

p prepare a dose, empty the packet contents into a container containing 2 tablespoons of ATER. DO NOT USE OTHER LIQUIDS OR FOODS.

clearance, this interaction is unlikely to be of clinical concern. Nonetheless, individual patients may require additional titration of their theophylline dosage when lansoprazole is started or stopped to ensure clinically effective blood levels.

In a study of healthy subjects neither the pharmacokinetics of warfarin enantiomers nor prothrombin time were affected following single or multiple 60 mg doses of lansoprazole. However, there have been reports of increased International Normalized Ratio (INR) and prothrombin time in patients receiving proton pump inhibitors, including lansoprazole, and warfarin concomitantly. Increases in INR and prothrombin time may lead to abnormal bleeding and even death. Pathents treated with proton pump inhibitors and warfarin concomitantly may need to be monitored for increases in INR and prothrombin time. Lansoprazole has also been shown to have no clinically significant interaction with amoxicillin. In a single-dose crossover study examining lansoprazole 30 mg and omeprazole 20 mg each administered alone and concomitantly with sucraflate. In refere, protron pump inhibitors should be taken at least 30 minutes prior to sucraflate, In clinical trials, antacids were administered acroace and sucrease and

50 mg/kg/day (13 times the recommended human dose based on body surface area) in a 1-year toxicity study.

In a 24-month carcinogenicity study, CD-1 mice were treated orally with doses of 15 to 600 mg/kg/day, 20 80 times the recommended human dose based on body surface area. Lansoprazole produced a dose-related increased incidence of gastric ECL cell hyperplasia. It also produced a dose-related increased incidence of surface in the surface area. Lansoprazole produced a dose-related increased incidence of liver tumors (hepatocellular adenoma plus carcinoma). The tumor incidences in male mice treated with 300 and 600 mg/kg/day (40 to 80 times the recommended human dose based on body surface area) and female intectreated with 150 to 600 mg/kg/day (20 to 80 times the recommended human dose based on body surface area) exceeded the ranges of background incidences in historical controls for this strain of mice. Lansoprazole treatment produced adenoma of rete testis in male mice receiving 75 to 600 mg/kg/day (10 to 80 times the recommended human dose based on body surface area).

Lansoprazole was not genotoxic in the Ames test, the ex vivo rat hepatocyte unscheduled DNA synthesis (IDS) test, the *in vivo* mouse micronucleus test or the rat bone marrow cell chromosomal aberration test. It was positive in *in vitro* human hymphocyte chromosomal aberration assays.

aberration assays.

Lansoprazole at oral doses up to 150 mg/kg/day (40 times the recommended human dose based on body surface area) was found to have no effect on fertility and reproductive performance of male and female rats.

Pregnancy: Teatogenic Effects.

Pregnancy Category B

Lansoprazole Teratology studies have been performed in pregnant rats at oral doses up to 150 mg/kg/day (40 times the recommended human dose based on body surface area) and pregnant rabbits at oral doses up to 30 mg/kg/day (16 times the recommended human dose based on body surface area) and have revealed no evidence of impaired fertility or harm to the fetus due to

lansoprazole.

There are, however, no adequate or well-controlled studies in pregnant women. Becauss animal reproduction studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed. Pregnancy Category C

See **WARNINGS** (above) and full prescribing information for clarithromycin before using in

ediatric Use
he safety and effectiveness of PREVACID have been established in pediatric patients 1 to
Paers of age for short-term treatment of symptomatic GERD and erosive esophagitis. Use
PREVACID in this population is supported by evidence from adequate and well-controlled
utilises of PREVACID in adults with additional clinical, pharmacokinetic, and
harmacodynamic studies performed in pediatric patients. The adverse events profile in
ediatric patients is similar to that of adults. There were no adverse events profile in
ediatric patients set were not previously observed in adults. The safety and effectiveness of
REVACID in patients <1 year of age have not been established.

PREVACID in patients <1 year of age have not been estaunished.

1 to 11 years of age
The pediatric safety of PREVACID Delayed-Release Capsules has been assessed in
66 pediatric patients aged 1 to 11 years of age. Of the 66 patients with GERD 85% (56/66)
took PREVACID for 8 weeks and 15% (10/66) took for 12 weeks.

The most frequently reported (2 or more patients) treatment-related adverse events in
patients to 11 years of age (N=66) were constipation (5%) and headache (3%).

seen in males. **Use in Geriatric Patients** Ulcer healing rates in elderly patients are similar to those in a younger age group. The

Itical (ridwide, over 10,000 patients have been treated with lansoprazole in Phase 2-3 clinical (ridwide, over 10,000 patients have been treated with lansoprazole in Phase 2-3 clinical (ridwide, over 10,000 patients) and output of treatment. The adverse reaction profiles for VACID Delayed-Release Oral Suspension similar. In general, lansoprazole treatment has been well-tolerated in both short-term long-term trials.

The following adverse events were reported by the treating physician to have a possible or bable relationship to drug in 1% or more of PREVACID-treated patients and occurred at reater rate in PREVACID-treated patients than placebo-treated patients and incidence of Possibly or Probably

Treatment-Related Adverse Events in Short-Term, Placebo-Controlled Studies

Treatment-Related Adverse Events in Short-Term, Placebo-Controlled Studies		
	PREVACID	Placebo
	(N= 2768)	(N= 1023)
lody System/Adverse Event	%	%
lody as a Whole		
Abdominal Pain	2.1	1.2
ligestive System		
Constipation	1.0	0.4
Diarrhea	3.8	2.3
Nausea	1.3	1.2
leadache was also seen at greater than 1% incidence but was more common on placeho		

The incidence of diarrhea was similar between patients who received placebo and patients who received lansoprazole 60 mg (2.9%, 1.4%, 4.2%, and 7.4%, respectively). The most commonly reported possibly or probably treatment-related adverse event during maintenance therapy was diarrhea. In the risk reduction study of PREVACID for NSAID-associated gastric ulcers, the incidence of diarrhea for patients treated with PREVACID was 5%, misoprostol 22%, and placebo 3%. Additional adverse experiences occurring in <a href="https://doi.org/10.1007/j.cm/10.1

voluntarily from a population of unknown size, estimates of frequency cannot be made. These events are listed below by QOSTART body system.

Body as a Whole- anaphylactoid-like reaction; Digestive System- hepatotoxicity, pancreatitis, vomiting, Hemic and Lymphatic System - agranulcytosis, aplastic anemia, hemolytic anemia, hemolytic and Lymphatic System - agranulcytosis, aplastic anemia, hemolytic anemia, leukopenia, neuropenia, pancytopenia, thrombocytopenia, and thrombotic thrombocytopenic purpura; Skin and Appendages – severe dermadologic reactions including erythema multiforme, Sevens-Johnson syndrome, toxic epidermal necrolysis (some fatal); Special Senses – speech disorder; Urogenital System – urinary retention.

Combination Therapy with Amostillia and Clarithromycin in clinical trials using combination therapy with PREVACID plus amoxicillin and clarithromycin, and PREVACID plus amoxicillin, no adverse reactions speculiar to these drug combinations were observed. Adverse reactions that have occurred have been limited to those that had been previously reported with PREVACID, amoxicillin, or clarithromycin. Triple Therapy: PREVACID/amoxicillin characteristics of the properties of the properties

formation on adverse reactions with amoxi rts, ADVERSE REACTIONS sections.

Laboratory Values The following changes in laboratory parameters for lansoprazole were reported as adverse

nts: normal liver function tests, increased SGOT (AST), increased SGOT (ALT), increased attinine, increased alkaline phosphatase, increased globulins, increased GGTP, eased/decreased/ahnormal MBC, abnormal AG ratio, abnormal ABC, bilirobinemia, inophilial, hyperlipiemia, increased/decreased/elcrotyles, increased/decreased/abnormal elesterol, increased glucocorticoids, increased LDH, increased/decreased/abnormal elests, and increased/agstrain levels. Unite abnormalities such as albumiumica, glycosumi, hematuria were also reported. Additional isolated laboratory abnormalities were rated.

and hematuria were also reported. Additional isolated lauviratory auroritarian reported. In the placebo controlled studies, when SGOT (AST) and SGPT (ALT) were evaluated, 0.4% (4978) placebo patients and 0.4% (11/2677) larsoprazole patients flad enzyme elevations greater than three times the upper limit of normal range at the final treatment vist. None of these lansoprazole patients reported jaundice at any time during the study. In clinical trials using combination therapy with PREVACID plus amoxicillin and clarithromycin, and PREVACID plus amoxicillin, no increased laboratory abnormalities particular to these drug combinations were observed.

For more information on laboratory value changes with amoxicillin or clarithromycin, refer to their package inserts, ADVERSE REACTIONS section.

to their package inserts, AUVENOR DEVENTIONS SEASON.

OVERDOSAGE

Oral doses up to 5000 mg/kg in rats (approximately 1300 times the recommended human dose based on body surface area) and nice (about 675.7 times the recommended human dose based on body surface area) did not produce deaths or any clinical signs.

Lanspirazole is not removed from the circulation by hemodialysis. In one reported case of overdose, the patient consumed 600 mg of lanspirazole with no adverse reaction.

Distributed by Distributed by Lake Forest, IL 60045, U.S.A.

ENSURE® is a registered trademark of Abbott Laboratories.

Del 12.5368.828 Rev. July 2004

RenOs-3586-R24 Rev. July, 2004
© 1995-2004 TAP Pharmaceutical Products Inc.
For more detailed information, see full prescribing information or contact TAP Medical Information at 1-800-622-2011.