PAP doesn’t cut rates of cardiovascular events, death

BY MARY ANN MOON
Frontline Medical News

Positive airway pressure, whether delivered continuously (CPAP) or as adaptive serv-ventilation, doesn’t reduce the rate of cardiovascular (CV) events or death in patients who have sleep apnea, according to a report published online July 11 in JAMA.

Positive airway pressure (PAP) relieves the symptoms of sleep apnea and has been reported to improve cardiovascular risk factors such as hypertension, insulin resistance, and endothelial dysfunction. However, whether the treatment improves “hard” vascular outcomes such as stroke and MI has never been established, said Jie Yu, MD, of the department of cardiology, Peking University and the Ministries of Health and Education, Beijing, and his associates.

They performed a systematic review of the literature and a meta-analysis of 10 randomized clinical trials that compared PAP against standard care or a sham treatment and had at least 6 months of follow-up for CV events. The meta-analysis involved 7,266 participants who had either obstructive (5,683 patients) or central (1,583 patients) sleep apnea. There were 356 major adverse CV events and 613 deaths during a median follow-up of 6-68 months.

The use of PAP showed no significant association with a range of outcomes: major adverse CV

Azithromycin improves QOL in asthmatics

Drug cut exacerbations

BY DOUG BRUNK
Frontline Medical News

Adults with persistent symptomatic asthma who took azithromycin as an add-on therapy experienced fewer exacerbations and had improved quality of life, compared with their peers who took a placebo, a multicenter, randomized trial demonstrated.

“Macrolide antibiotics have antibacterial, anti-viral, and anti-inflammatory effects, and are reported to be beneficial in both eosinophilic and noneosinophilic subtypes,” a group of Australian researchers wrote online July 4 in The Lancet (doi: org/10.1016/S0140-6736[17]31281-3). “Systematic reviews of randomized, controlled trials report benefits of macrolides on asthma symptoms but [we] are unable to draw conclusions about the effects on other endpoints, including exacerbations, due to lack of data, heterogeneity of results, and inadequate study design and sample size.”

Led by Peter G. Gibson, MBBS, of Hunter Medical Research Institute, New South Wales, Australia, researchers at eight clinical sites con-

ADD-ON THERAPY CUTS EXACERBATIONS

continued on page 4

Reasons for inpatient stays shift

The reasons for inpatient stays have shifted. See 2014’s most frequent diagnoses for hospital stays, following pregnancy/childbirth and newborns and neonates, on page 7.
ducted a randomized trial to test the hypothesis that the macrolide antibiotic azithromycin reduces asthma exacerbations and improves quality of life in patients with symptomatic asthma on inhaled maintenance therapy. To be eligible for the trial, known as Asthma and Macrolides: the Azithromycin Efficacy and Safety Study, or AMAZES, patients had to be at least 18 years of age, be using an inhaled corticosteroid and long-acting bronchodilator, and have no hearing impairment or abnormal prolongation of the corrected QT interval. Primary efficacy endpoints were the total number of asthma exacerbations (severe and moderate) over 48 weeks and asthma quality of life based on responses to the Asthma Quality of Life Questionnaire (Chest. 1999 May;115[5]:1265-70). Of the 420 patients, 213 were allocated to take 500 mg azithromycin three times weekly and 207 were allocated to placebo. In all, 168 patients in the azithromycin group completed 48 weeks of treatment, compared with 166 in the placebo group. Their median age was 60 years, 76% had

Add-on therapy cuts exacerbations // continued from page 1

Continued on page 7
The impact on community microbial resistance remains unclear

Since microbial resistance is a well-known side effect of antibiotic use, add-on therapy with azithromycin in asthma needs to be restricted to those patients with the highest unmet medical need (for example, frequent exacerbators) and to time periods with the greatest risk of exacerbations (such as winter). Biomarkers that predict the therapeutic response to macrolides might facilitate optimal patient selection. Further research is needed to elucidate the most important mechanism of action of these pleiotropic drugs. Macrolides have anti-inflammatory, antibacterial, and antiviral effects. However, the authors did not observe a reduction in inflammatory cell counts in sputum to support a definite anti-inflammatory effect. Azithromycin also was effective in patients with and without potentially pathogenic microorganisms in sputum cultures at baseline. Since azithromycin reduced both asthma exacerbations and respiratory infections, the benefits of azithromycin might be caused by preventing viral-induced attacks in asthma. Azithromycin stimulates phagocytosis of microbes and dead cells by macrophages (i.e., efferocytosis), an effect that is likely to be independent of the nature of the accompanying neutrophilic or eosinophilic airway inflammation.

Dr. Gibson and his colleagues have clearly shown that add-on therapy with azithromycin is effective and safe in adult patients with uncontrolled asthma despite treatment with inhaled corticosteroids and long-acting beta-agonists. Azithromycin benefited patients with both eosinophilic and noneosinophilic asthma. However, the effects of long-term therapy with macrolides on community microbial resistance remain a public health concern. Future studies with potentially safer nonantibiotic macrolides in uncontrolled severe asthma are warranted. Since the antimicrobial effects probably contribute to the overall efficacy of macrolides, the beneficial effects of nonantibiotic macrolides might be intermediate between macrolide antibiotics and placebo.

This text is excerpted from a commentary published online July 4 in The Lancet (doi.org/10.1016/S0140-6736[17]31547-7). Guy Brusselle, MD, is with the department of respiratory medicine at Ghent (Belgium) University Hospital and Ian Pavord, MD, is with the University of Oxford’s Nuffield Department of Medicine, in England. Both authors disclosed having received honoraria and other financial support from numerous pharmaceutical companies.
**NEWS**

**Frequent bronchiectasis exacerbations linked to higher mortality**

**BY MITCHEL L. ZOLER**
Frontline Medical News

WASHINGTON – Bronchiectasis patients with three or more exacerbations per year had twice the mortality during 5-year follow-up as patients with no recent exacerbations, in a prospective registry of nearly 2,600 European bronchiectasis patients.

A multivariate analysis showed this statistically significant doubled death rate after adjustment for baseline demographic and clinical differences between patients with no exacerbations during the year before they entered the registry, James D. Chalmers, MD, said at an international conference of the American Thoracic Society.

Having had frequent exacerbations at a rate of three or more annually prior to enrollment was common, with 37% of the 2,596 bronchiectasis patients in the registry having this history, said Dr. Chalmers, a pulmonologist at the University of Dundee, Scotland. This 37% prevalence contrasted with a 19% U.S. prevalence of bronchiectasis patients having two or more exacerbations per year among 2,114 patients enrolled in a 13-center U.S. registry that was reported during the same session by Timothy R. Aksamit, MD, a pulmonologist at the Mayo Clinic in Rochester, Minn. Dr. Aksamit contended that the U.S. registry tried to exclusively enroll patients with bronchiectasis and no other disorder, possibly explaining the prevalence difference between Europe and the United States.

The European registry included patients with bronchiectasis seen in 10 centers in seven European countries and Israel. They averaged 67 years of age. While more than a third had a history of at least three exacerbations a year, one-quarter had no exacerbations during the year before they entered the study.

The prospective study also showed that, among patients with three or more exacerbations annually, the risk for a subsequent exacerbation was five times higher than among patients with no recent exacerbations.

The U.S. registry reported by Dr. Aksamit had two-year follow-up data for 1,049 of the enrolled patients, a subgroup that closely matched the entire population initially enrolled. The 2-year follow-up showed an overall average exacerbation rate of 0.75 episodes per year, but this was driven largely by the subgroup of patients who entered the registry with a history of two or more exacerbations per year, who then averaged about 2.6 exacerbations during follow-up. In contrast, patients who entered the registry with a history of fewer than two exacerbations per year averaged fewer than a third of an exacerbation per year during follow-up.

The European bronchiectasis registry was partially funded by Bayer. Dr. Chalmers has been a consultant to Bayer and to AstraZeneca, Basilea, Grifols, Napp, and Raptor and has received research funding from Aradigm, AstraZeneca, Bayer, GlaxoSmithKline, and Pfizer.

[Dr. Chalmers and Dr. Aksamit]

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**View on the News**

Eric Gartman, MD, FCCP, comments: The longitudinal outcomes of patients within both of these bronchiectasis cohorts demonstrate that there is a subset of patients with this condition who are prone to exacerbation and that this high exacerbation rate portends a very poor prognosis. As such, increased focus should be placed on this particular group of patients in an attempt to prevent exacerbations and subsequent clinical decline.

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**News From Chest**

**CRITICAL CARE COMMENTARY**

Medical students: The role of conscience in medicine.

Vera A. De Palo, MD, MBA, FCCP, is Medical Editor in Chief of CHEST Physician.

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Septicemia admissions almost tripled from 2005 to 2014

BY RICHARD FRANKI
Frontline Medical News

Admissions for septicemia nearly tripled from 2005 to 2014, as it became the third most common diagnosis for hospital stays, according to the Agency for Healthcare Research and Quality. There were more than 1.5 million hospital stays with a principal diagnosis of septicemia in 2014, an increase of 192% over the 518,000 stays in 2005. The only diagnoses with more admissions in 2014 were pregnancy/childbirth with almost 4 million stays and newborns/neonates at almost 4 million, although both were down from 2005. That year, septicemia did not even rank among the top 10 diagnoses, the AHRQ reported.

Osteoarthritis was the fourth most common diagnosis in 2014 with almost 1.1 million stays, an increase of almost 50% from 2004, when it was the seventh most common diagnosis. Admissions for the fifth most common diagnosis in 2014, congestive heart failure, were down by over 14% from 2005, data from the National Inpatient Sample show. Pneumonia, which was the third most common diagnosis in 2005, dropped by 32% and ended up in sixth place in 2014, while admissions for coronary atherosclerosis, which was fourth in 2005, decreased by 63%, dropping out of the top 10 by 2014, the AHRQ said.

Septicemia was the most common diagnosis for inpatient stays among those aged 75 years and older and the second most common for those aged 65-74 and 45-64. The leading nonmaternal, non-neonatal diagnosis in the two youngest age groups, 0-17 and 18-44 years, was mood disorders, and the most common cause of admissions for those aged 45-64 and 65-74 years was osteoarthritis, the AHRQ reported.

PAP didn’t improve blood pressure // continued from page 1

...clinical trials] suggests that the association [between] sleep apnea and vascular disease risk may represent disease processes that cannot be ameliorated by PAP delivered at the average intensity..."

The evidence from these randomized clinical trials] suggests that the association [between] sleep apnea and vascular..."

Daniel J. Gottlieb, MD, is in the medical service at the VA Boston Healthcare System and in the division of sleep medicine at Harvard Medical School, Boston. He reported receiving personal fees from VIVUS. Dr. Gottlieb made these remarks in an editorial accompanying Dr. Yu’s report (JAMA. 2017;318:128-30).
Tool predicts antimicrobial resistance in sepsis

BY HEIDI SPLETE
Frontline Medical News

Use of a clinical decision tree predicted antibiotic resistance in sepsis patients infected with gram-negative bacteria, based on data from 1,618 patients. Increasing rates of bacterial resistance have “contributed to the unwarranted empiric administration of broad-spectrum antibiotics, further promoting resistance emergence across microbial species,” said M. Cristina Vazquez Guillelmet, MD, of the University of New Mexico, Albuquerque, and her colleagues (Clin Infect Dis. 2017 Jul 10. doi: 10.1093/cid/cix612).

The researchers identified adults with sepsis or septic shock caused by bloodstream infections who were treated at a single center between 2008 and 2015. They developed clinical decision trees using the CHAID algorithm (Chi-squared Automatic Interaction Detection) to analyze risk factors for resistance associated with three antibiotics: piperacillin-tazobactam (PT), cefepime (CE), and meropenem (ME).

“We found good overall agreement between the accuracies of the [multivariable logistic regression] models and the decision tree analyses for predicting antibiotic resistance,” the researchers said.

Overall, resistance rates to PT, CE, and ME were 29%, 22%, and 9%, respectively, and 6.6% of the isolates were resistant to all three antibiotics. Factors associated with increased resistance risk included residence in a nursing home, transfer from an outside hospital, and prior antibiotics use. Resistance to ME was associated with infection with Pseudomonas or Acinetobacter spp, the researchers noted, and resistance to PT was associated with central nervous system and central venous catheter infections.

Clinical decision trees were able to separate patients at low risk for resistance to PT and CE, as well as those with a risk greater than 30% of resistance to PT, CE, or ME. “We also found good overall agreement between the accuracies of the [multivariable logistic regression] models and the decision tree analyses for predicting antibiotic resistance,” the researchers said.

The findings were limited by several factors, including the use of data from a single center and incomplete reporting of previous antibiotic exposure, the researchers noted. However, the results “provide a framework for how empiric antibiotics can be tailored according to decision tree patient clusters,” they said.

Ribaxamase prevented C. difficile infections by protecting microbiome

BY MICHELE G. SULLIVAN
Frontline Medical News

VIENNA – An investigational beta-lactamase reduced Clostridium difficile infections by 71% in patients receiving extended antibiotic therapy for respiratory infections but not by killing the opportunistic bacteria.

Rather, ribaxamase prevented C. difficile infections (CDI) by breaking down excess therapeutic antibiotics in the gut before they could injure an otherwise healthy microbiome, John Kokai-Kun, PhD, said at the European Society of Clinical Microbiology and Infectious Diseases annual congress.

“Up to 50% of an antibiotic dose is excreted into the small intestine, where it starts to disrupt the bowel microbiome and predisposes you to pick up C. difficile,” said Dr. Kokai-Kun, vice president of non-clinical affairs at Synthetic Biologics, Rockville, Md. “Ribaxamase is designed to block this cascade. If we protect the microbiome, any C. difficile that finds its way in would not find a gut conducive to the germination of vegetative cells.”

Ribaxamase is an oral enzyme that breaks the lactam ring in penicillins and cephalosporins. It’s formulated to release at a pH of 5.5 or higher, an environment that begins to develop in the upper small intestine near the bile duct – the same place that excess antibiotics are excreted.

“The drug is intended to be administered during, and for a short time after, intravenous administration of specific beta-lactam-containing antibiotics,” Dr. Kokai-Kun said. Ribaxamase doesn’t work on carbapenem-type antibiotics, he noted, and Synthetic Biologics is working on an effective enzyme for those as well.

In early human studies, ribaxamase was well tolerated and didn’t interfere with the pharmacokinetics of therapeutic antibiotics (Antimicrob Agents Chemother. 2017 Mar;61[3]:e02197-16). It’s also effective in patients who are taking a proton pump inhibitor, he said.

Dr. Kokai-Kun reported the results of a phase 2b study of 412 patients who received IV ceftriaxone for lower respiratory infections. They were assigned 1:1 to either 150 mg ribaxamase daily or placebo throughout the IV treatment and for 3 days after.

The primary endpoint was prevention of C. difficile infection. The secondary endpoint was prevention of non-C. difficile antibiotic-associated diarrhea. An exploratory endpoint examined the drug’s ability to protect the microbiome. Patients were monitored for 6 weeks after treatment stopped.

The cohort was a mean 70 years old. One-third of patients also received a macrolide during their hospitalization, and one-third were taking proton pump inhibitors. The respiratory infection cure rate was about 99% in both groups at both 72 hours and 4 weeks.

Eight patients in the placebo group (3.8%) and two in the active group (less than 1%) developed C. difficile infection. That translated to a statistically significant 71% risk reduction, with a P value of .027, Dr. Kokai-Kun said. Ribaxamase did not hit its secondary endpoint of preventing all-cause diarrhea or antibiotic-associated diarrhea that was not caused by C. difficile infection.

Although not a primary finding, ribaxamase also inhibited colonization by vancomycin-resistant enterococci, which occurred in about 70 (40%) patients in the placebo group and 40 (20%) in the ribaxamase group at both 72 hours and 4 weeks.

All patients contributed stool samples at baseline and after treatment for microbiome analysis. That portion of the study is still ongoing, Dr. Kokai-Kun said.

Synthetic Biologics sponsored the study and is developing ribaxamase. Dr. Kokai-Kun is the company’s vice president of nonclinical affairs.
CRITICAL CARE MEDICINE

AR-301 holds promise for S. aureus pneumonia

BY DAMIAN MCNAMARA
Frontline Medical News

NEW ORLEANS – Monoclonal antibody therapies have already upended treatment strategies in cancer, dermatology, and multiple inflammatory diseases, and infectious disease may be next.

That’s because a single injection of a monoclonal antibody in development, AR-301, appeared to be safe and effective as an adjunct treatment for severe pneumonia caused by Staphylococcus aureus, according to a new study. The monoclonal antibody attacks the alpha-toxin secreted by S. aureus, thereby helping to protect immune cells.

Researchers assessed 48 patients between May 2012 and May 2016 in a randomized, double-blind, placebo-controlled trial. Each participant received a single injection of placebo or AR-301 (at one of four doses) to test the antibody’s tolerability and effectiveness.

“We know S. aureus pneumonia is a big problem. There is a lot of antibiotic resistance, and that is why we need new treatments,” Celine Gonzalez, MD, of the Dupuytren Central University Hospital in Limoges, France, said in an interview.

“Animal studies have shown the monoclonal antibody seems to be useful. This is the first in-human study to use a monoclonal antibody to treat hospital-acquired pneumonia due to Staphylococcus aureus,” Dr. Gonzalez said in a late-breaking poster presentation at the annual meeting of the American Society for Microbiology.

Treatment started within 36 hours of onset of severe pneumonia. Severity was based on a mean PaO$_2$/FiO$_2$ of 147 and/or a need for catecholamine. Six cases of pneumonia were related to MRSA and the remaining 42 to methicillin-susceptible S. aureus. The mean APACHE II score was 18.7, the mean Clinical Pulmonary Infection Score was 9.6, and the mean Sequential Organ Failure Assessment score was 6.9.

Participants were recruited from

Continued on following page

Dr. Celine Gonzalez

Damian McNamara/Frontline Medical News
13 ICUs in four countries. About 80% of participants were men. Their mean age was 56 years, and mean body mass index was 29 kg/m². Concurrent antibiotic treatment choice and duration were at the investigator’s discretion.

*S. aureus* infection was considered eradicated if a follow-up culture was negative, a result achieved by 63% of the 16 placebo patients and 75%-88% of the AR-301-dosage groups. Eradication was also based on observed clinical success in the absence of a confirmatory culture. This was achieved by 38% in the placebo group and 13%-25% of the monoclonal antibody cohorts. A total of seven placebo patients and 15 AR-301 patients met eradication by these criteria.

Side effects were primarily minor and transient, Dr. Gonzalez said. Of the 343 total adverse events reported, only 8 (2.3%) were considered treatment related, she added.

“Animal studies have shown the monoclonal antibody seems to be useful. This is the first in-human study to use a monoclonal antibody to treat hospital-acquired pneumonia due to *Staphylococcus aureus*,” Dr. Celine Gonzalez said.
“In infectious disease, it’s the beginning” for monoclonal antibody therapy, Dr. Gonzalez said. “But, it appears to be the future because … it is a more specific treatment, and there is no resistance.”

The study suggests adjunctive treatment with AR-301 appears safe for treatment of hospital-acquired bacterial pneumonia, she noted.

The next step will be to confirm the findings in a larger, follow-up study that includes more efficacy outcomes, Dr. Gonzalez added.

Dr. Gonzalez reported having no relevant disclosures. The study’s principle investigator is a scientific advisor for Aridis Pharmaceuticals, which is developing AR-301.

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A combination of ambrisentan (Letairis) and tadalafil (Cialis) improves regional and global right ventricular contractility in patients with scleroderma-associated pulmonary arterial hypertension, according to an open-label investigation of 23 patients. The project was a follow-up to a previous report showing that the upfront combination – tadalafil 40 mg and ambrisentan 10 mg oral once daily – improved hemodynamics, right ventricular (RV) structure and function, and functional status in treatment-naive patients after 36 weeks and "may represent a very effective therapy for this patient population" (Am J Respir Crit Care Med. 2015 Nov 1;192[9]:1102-10).
Survival in scleroderma pulmonary arterial hypertension (PAH) depends mostly on RV function, so investigators in the follow-up study wanted to take a closer look at how the combination affected the heart. They reviewed conventional echocardiograph imaging and RV strain analyses for the 23 of the 24 patients in the original trial for which it was available (Am J Respir Crit Care Med. 2017 Jun 29. doi: 10.1164/rcm.201704-0789LE).

At baseline, the subjects had normal left ventricular (LV) size and function, with borderline left atrial enlargement and mild LV diastolic dysfunction. Their right heart chambers were significantly dilated, with RV hypertrophy. Conventional RV function parameters—tricuspid annular systolic plane excursion (TAPSE) and fractional area change (FAC)—were impaired. RV systolic pressure (RVSP) was severely elevated. There was also a marked reduction of global RV longitudinal systolic strain (RVLSS), compared with normal values, mainly because of a reduction in midventricular and apical RVLSS, with relative hyperkinesis of basal RVLSS.

Continued on following page
Amplatzer devices outperform oral anticoagulation

BY BRUCE JANCIN
Frontline Medical News

PARIS – Percutaneous left atrial appendage closure with an Amplatzer device in patients with nonvalvular atrial fibrillation was associated with significantly lower rates of all-cause and cardiovascular mortality, compared with oral anticoagulation, in a large propensity score–matched observational registry study. Left atrial appendage closure (LAAC) also bested oral anticoagulation (OAC) with warfarin or a novel oral anticoagulant (NOAC) in terms of net clinical benefit on the basis of the device therapy’s greater protection against stroke and systemic embolism coupled with a trend, albeit not statistically significant, for fewer bleeding events, Steffen Gloekler, MD, reported at the annual congress of the European Association of Percutaneous
Cardiovascular Interventions.

The Watchman LAAC device, commercially available both in Europe and the United States, has previously been shown to be superior to OAC in terms of efficacy and noninferiority regarding safety. But there have been no randomized trials of an Amplatzer device versus OAC. This lack of data was the impetus for Dr. Gloekler and his coinvestigators to create a meticulously propensity-matched observational registry.

Five hundred consecutive patients with AF who received an Amplatzer Cardiac Plug or its second-generation version, the Amplatzer Amulet, during 2009-2014 were tightly matched to an equal number of AF patients on OAC based on age, sex, body mass index, left ventricular ejection fraction, renal function, coronary artery disease status, hemoglobin level, CHA2DS2-VASc score, and HAS-BLED score. During a mean 2.7 years, or 2,645 patient-years, of follow-up, the composite primary efficacy endpoint, composed of stroke, systemic embolism, and cardiovascular or unexplained death occurred in 5.6% of the LAAC group, compared with 7.8% of controls in the OAC arm, for a statistically significant 30% relative risk reduction. Disabling stroke occurred in 0.7% of Amplatzer patients versus 1.5% of controls. The ischemic stroke rate was 1.5% in the device therapy group and 2% in the OAC arm.

All-cause mortality occurred in 8.3% of Amplatzer patients and 11.6% of the OAC group, for a 28% relative risk reduction. The cardiovascular death rate was 4% in the Amplatzer group, compared with 6.5% of controls, for a 36% risk reduction.

The composite safety endpoint, comprising all major procedural adverse events and major or life-threatening bleeding during follow-up, occurred in 3.6% of the Amplatzer group and 4.6% of the OAC group, for a 20% relative risk reduction that is not significant at this point because of the low number of events. Major, life-threatening, or fatal bleeding occurred in 2% of Amplatzer recipients versus 5.5% of controls, added Dr. Gloekler of University Hospital in Bern, Switzerland.

The net clinical benefit, a composite of death, bleeding, or stroke, occurred in 8.1% of the Amplatzer group, compared with 10.9% of controls, for a significant 24% reduction in relative risk in favor of device therapy.

Of note, at 2.7 years of follow-up only 55% of the OAC group were on warfarin, and 17% were taking a NOAC. At that point, 8% of the Amplatzer group were on any anticoagulation therapy.

Discussion of the study focused on that low rate of medication adherence in the OAC arm. Dr. Gloekler’s response was that, after looking at the literature, he was no longer surprised by the finding that only 55% of the control group were on OAC at follow-up.

“If you look in the literature, that’s exactly the real-world adherence for OACs. Even in all four certification trials for the NOACs, the rate of discontinuation was 30% after 2 years – and these were controlled studies. Ours was observational, and it depicts a good deal of the problem with any OAC in my eyes,” Dr. Gloekler said.

Patients on warfarin in the real-world Amplatzer registry study spent on average a mere 30% of time in the therapeutic international normalized ratio range of 2-3.

Dr. Gloekler reported receiving research funds for the registry from the Swiss Heart Foundation and Abbott.

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Biomarker distinguishes ARDS, acute heart failure

BY MITCHEL L. ZOLER
Frontline Medical News

WASHINGTON — Plasma levels of an interleukin-33 receptor that’s involved in inflammation regulation appeared able to discriminate between acute respiratory distress syndrome and acute uncompensated heart failure in an analysis with 72 patients.

In a second study, high plasma levels of the same interleukin-33 receptor, soluble suppressor of tumorigenicity 2 (sST2), identified acute respiratory distress syndrome (ARDS) patients who were sicker and more responsive to conservative fluid management, Sean D. Levy, MD, said at an international conference of the American Thoracic Society.

While further validation of sST2 is needed, its future as a clinically useful biomarker also depends on development of a test that could be easily and repeatedly used at the bedside, said Dr. Levy, a pulmonologist at New England Deaconess Medical Center in Boston. “We’re not quite there yet,” he explained. The sST2 test he used for his studies is sold by Critical Diagnostics.

In order to assess the ability of sST2 to reliably distinguish patients with ARDS from those with acute uncompensated heart failure, he and his associates selected 72 patients seen at the Massachusetts General Hospital in Boston with an initial diagnosis of acute uncompensated heart failure accompanied by bilateral lung infiltrates and acute hypoxemia requiring endotracheal intubation and mechanical ventilation. The investigators measured the sST2 level in a plasma specimen from each patient. In addition, after each patient either left the hospital or died, their case underwent review by two critical care physicians who retrospectively rediagnosed the patients as either having ARDS or acute uncompensated heart failure. This divided the cohort into 30 patients with ARDS and 42 with true acute heart failure. The two subgroups matched up fairly closely for most clinical measures and comorbidities, but APACHE III (Acute Physiology and Chronic Health Evaluation III) scores averaged significantly higher in the ARDS patients.

The plasma levels of sST2 showed a dramatic split between the two subgroups. The 30 patients retrospectively diagnosed with ARDS had an average level of 386 ng/mL with an interquartile range of 318–611 ng/mL. The 42 acute uncompensated heart failure patients averaged a sST2 level of 148 ng/mL, with an interquartile range of 84–225 ng/mL. The area under the receiver operator curve for discriminating between ARDS and acute heart failure using a cutpoint of 271 mg/mL was 0.86, showing “good” discrimination, Dr. Levy said. This cutpoint had a sensitivity of 83% and specificity of 88% for correctly distinguishing between ARDS and acute heart failure.

In a second analysis, Dr. Levy and his associates looked at the ability of sST2 levels to separate out patients with acute lung injury who had a more robust response to either the conservative or liberal fluid-management strategies tested in the Fluid and Catheter Treatment Trial (FACTT), run by the National Heart, Lung, and Blood Institute’s ARDS Clinical Trials Network. The primary outcome of FACTT was death from any cause 60 days after entry, and this showed no significant difference between conservative (restricted fluids and increased urine output) and liberal (the reverse) fluid management strategies in acute lung injury patients (N Engl J Med. 2006 Jun 15;354[14]:2564-75). From among the 1,001 patients enrolled in FACTT, 826 had specimens available for measuring sST2 (Crit Care Med. 2013 Nov;41[11]:2521-31). The researchers applied the sST2 cut point they derived in the first analysis to the FACTT cohort and identified 133 (16%) patients with a low sST2 level and 693 (84%) with a high level. The patients with high sST2 had significantly higher APACHE III scores, worse acidemia, and worse renal function.

Patients with high sST2 levels had a significant increase in ventilator-free days on conservative fluid management, compared with liberal management, while the two management strategies produced virtually identical results in the patients with low levels of sST2. Patients with high sST2 also had a significantly quicker time to extubation on a conservative strategy, compared with the liberal strategy, and again this correlation did not exist among patients with low sST2. However, as in the overall trial, a conservative strategy had no discernible impact on 60-day mortality, compared with the liberal strategy, even in the subgroup with high sST2.

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More pulmonary patients getting palliative care

BY RICHARD FRANKI
Frontline Medical News

Patients referred to palliative care are most likely to have cancer, but the proportion has gone down since 2009 as other diagnoses have increased, according to a report from the National Palliative Care Registry. In 2015, cancer patients made up 26% of the patients referred to palliative care, compared with 35% in 2009. The situation was reversed for the next three most common diagnoses in 2015: Cardiac diagnoses rose from 5% in 2009 to 13%, pulmonary diagnoses increased from 6% to 12%, and neurologic diagnoses went from 3% to 8%, the report showed.

Referrals by specialty were led by hospital medicine, which accounted for 48% of all patients referred to palliative care in 2015, with internal medicine/family medicine next at 14%, followed by pulmonary/critical care at 13% and oncology at 7%.

An increase in overall palliative care penetration was seen from 2009 to 2015, as the percentage of annual hospital admissions seen by a palliative care team increased from 2.7% to 4.8%. Over that same period, the percentage of palliative care patients who died in the hospital decreased from 29% to 22%, according to the report.

In 2015, there were 420 palliative care programs participating in the registry, which is a joint project of the Center to Advance Palliative Care and the National Palliative Care Research Center.

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VIEW ON THE NEWS
Daniel Ouellette, MD, FCPP, comments: “But doc, isn’t hospice just for cancer patients?” My 80-year-old patient has COPD, requires oxygen at 4 L/min at rest, and cannot walk to his mailbox despite being on a maximum bronchodilator regimen. Too old to be a candidate for lung transplant, I have few additional medical treatments to offer him. I hope that I can help him have comfort during the last days of his life. My response to him is: “Not any longer.” The study from the National Palliative Care Registry demonstrates that pulmonary physicians and their patients are increasingly aware that palliation plays an important role in the management of patients with end-stage respiratory disease.
Patients report issues with home O₂

BY KATIE WAGNER LENNON
Frontline Medical News

WASHINGTON – Patient education in the use of home oxygen halves the number of system use issues reported by patients, based on results of a survey of nearly 2,000 patients.

Pulmonary clinicians and patients report “intolerable barriers to home oxygen services,” lead researcher Susan S. Jacobs, RN, MS, said in a poster session at an international conference of the American Thoracic Society. These barriers include insufficient oxygen supply, inadequate and physically unmanageable portable options, and equipment malfunction.

In their study, Ms. Jacobs and her colleagues sought to determine the frequency and types of problems experienced by adult home oxygen users in the United States. Survey respondents were recruited via efforts by the ATS Public Advisory Roundtable. Links to the survey were posted on various patient advocacy websites, and flyers were posted at clinics and pulmonary rehabilitation programs asking patients to participate in an online, 60-item survey developed by the ATS Nursing Oxygen Working Group. Participants included 1,926 patients, but not all patients responded to every question.

“We’ve demonstrated that, if the patients are educated by a health-care professional, the problems with oxygen go down, said Susan Jacobs, who is a nurse coordinator in the division of pulmonary and critical care medicine at Stanford University.

“Most of the reported problems were related to respondents not having portable systems that let them be out of their house for more than 2-4 hours or [to systems that] were too heavy for the patients to lift up and down their stairs and out of their cars, and they had problems operating them,” Ms. Jacobs said.

The survey respondents also reported experiencing delivery problems, not being able to change the company providing them with oxygen, receiving incorrect or delayed orders from a physician, or being unable to get liquid oxygen. These responses were provided by 267, 177, 166, and 68 patients, respectively.

“Their work would seem to indicate that a physician can dictate exactly what system they want. ... You can try to give [patients] a lighter system, a backpack, a smaller tank, more tanks per week, depending on their lifestyle and their needs. But physicians, a lot of times, like all of us and our patients, [are] not aware of all these choices,” she said during the interview.

An online resource providing all of the pros and cons of the different types of portable oxygen systems that would be appropriate for physicians, nurses, and patients, as well as an examination of the quality standards of the oxygen suppliers, are needed, she noted.

Ms. Jacobs reported no financial disclosures.

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VIEW ON THE NEWS
Vera A. De Palo, MD, MBA, FCCP, comments: The authors point out that there are a multitude of reasons that a patient may have difficulty with oxygen therapy. Their work would seem to indicate that conversation between the care team members (patient/family, physician, and respiratory therapy provider) can help reduce the questions and difficulties that a patient and his/her family may have after the prescribed therapy has been delivered. Any action that would enhance the likelihood of compliance with the prescribed therapy would be a benefit to our patients.
Algorithm for identifying IPF has low PPV

BY M. ALEXANDER OTTO
Frontline Medical News

ICD-9 codes were poor at picking out idiopathic pulmonary fibrosis patients from administrative databases for epidemiologic studies, but a new tool could improve diagnostic accuracy, according to Kaiser Permanente and University of California, San Francisco, investigators.

“In the age of large administrative databases and electronic medical records, there is rich opportunity to conduct population-based studies of disease behavior, outcomes, health care use, and other matters, but researchers first need to be able to accurately identify patients with idiopathic pulmonary fibrosis (IPF) in large data sets, said investigators led by Brett Ley, MD, an assistant professor of medicine at UCSF.

The research community has traditionally relied on claims for specific IPF diagnostic codes – ICD-9 code 516.3 or ICD-9-CM code 516.31 – to identify patients, but the approach had never been validated. To see how well it works, the investigators applied it to the nearly 5.4 million adults in the Kaiser Permanente Northern California system during 2000-2014. After patients with interstitial lung disease-associated codes entered on or after the day of the last IPF code were excluded, the algorithm identified 2,608 patients as having IPF (Ann Am Thorac Soc. 2017 Jun;14[6]:880-7). Next, the investigators randomly

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Evans R. Fernandez Perez, MD, is a pulmonologist at National Jewish Health, Denver. He made his comments in an editorial, and reported speaker’s fees from Boehringer Ingelheim and Genentech (Ann Am Thorac Soc. 2017 Jun;14[6]:829-30).
selected 150 of those patients and examined their medical records, procedure codes, CTs, and other patient-level data to see how many of them really had IPF. The results weren’t good. The positive predictive value of the IPF code-based algorithm was only 42.2%, with a sensitivity of 55.6%.

The widely used code-based IPF algorithm does “not generate accurate estimates of IPF incidence and prevalence. ... Over half of the patients identified as having IPF ... did not have IPF on case review. Alarmingly, whereas half of the misclassified cases had an alternative [interstitial lung disease] diagnosis, the other half had no clinical or radiologic evidence of ILD [interstitial lung disease] at all.” The algorithm also “likely misses a substantial proportion of patients who do have IPF,” Dr. Ley and his colleagues said. “We can only speculate about the reasons. ... It seems likely to be due to a combination of misdiagnosis at the clinical level and miscoding at the administrative level,” they said.

To try to improve the situation, the team tweaked the algorithm to include only patients 50 years or older who had at least two 516.3 or 516.31 claims 1 month or more apart and a chest CT procedure code beforehand. They again excluded ILD-associated claims on or after the day of the last IPF code. Although the sensitivity of the algorithm does “not generate accurate estimates of IPF incidence and prevalence. ... Over half of the patients identified as having IPF ... did not have IPF on case review. Alarmingly, whereas half of the misclassified cases had an alternative [interstitial lung disease] diagnosis, the other half had no clinical or radiologic evidence of ILD [interstitial lung disease] at all.” The algorithm also “likely misses a substantial proportion of patients who do have IPF,” Dr. Ley and his colleagues said. “We can only speculate about the reasons. ... It seems likely to be due to a combination of misdiagnosis at the clinical level and miscoding at the administrative level,” they said.

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modified algorithm was lower than the original, it had a more robust positive predictive value of 70.4% in the derivation cohort and 61.8% in the validation cohort, both derived from the 150 patients used to validate the original algorithm.

“By making a few simple, empirically derived changes to the IPF algorithm,” it’s possible to “more reliably [identify] patients” with IPF. “We believe the modified IPF algorithm will be useful for population-based studies of IPF … that require high diagnostic certainty,” the investigators concluded.

The traditional algorithm found an incidence of 6.8 cases per 100,000 person-years, which was on the low end of previous reports, perhaps because of the relative health and youth of the 5.4 million patient pool. As in past studies, IPF incidence increased with older age and was highest in white patients and men.

The researchers called for further study of whether the more specific codes will allow for improved case classification of IPF.

The work was funded by the National Institutes of Health. Dr. Ley reported speaker’s fees from Genentech, and another author was an employee of Genentech. The senior author Harold Collard, MD, an associate professor in UCSF’s division of pulmonary and critical care medicine, reported personal fees from various companies.

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Continued from previous page
GI disorder risk may rise in poorly controlled asthma

BY BRIAN HOYLE
Frontline Medical News

SAN FRANCISCO – Pediatric patients who have asthma that is poorly controlled may be more likely to have functional gastrointestinal (GI) disorders, which feature chronic GI distress that has several causes, according to a study of patients treated at one hospital. Female sex and increased anxiety were influential factors.

“This study suggests a high prevalence of functional GI disorders among patients with persistent asthma. Moreover, patients with functional GI disorders had poor asthma control and increased anxiety. Clinicians should consider functional GI disorders in patients with poor asthma control and assess for anxiety as indicated,” Ruben J. Colman, MD, a pediatric resident at SBH Health System, New York, said at the Pediatric Academic Societies meeting.

The prospective, cross-sectional study recruited patients aged 4-20 years at the emergency department, pediatric inpatient unit, and ambulatory clinics at St. Barnabas Hospital, a 422-bed, not-for-profit, acute care community hospital. Those with persistent asthma, which was evident by an ongoing history of daily inhaled corticosteroid medication, were enrolled.

Functional GI disorders including functional abdominal pain, irritable bowel syndrome, and functional dyspepsia were evaluated. The study was prompted by the knowledge that these conditions are a common cause of chronic GI symptoms in children, and from the findings of a retrospective study of 30,000 patients in Europe that reported a higher prevalence of asthma in those with functional GI disorders, compared with those without chronic GI distress (Aliment Pharmacol Ther. 2014 Aug;40[40]:382-91). Data are scarce in North America concerning asthma control and functional GI disorders in both pediatric and adult populations.

The validated Questionnaire on Pediatric Gastrointestinal Symptoms–Rome III version was used to assess functional GI disorders. Asthma control was assessed using the childhood Asthma Control Test.

Continued on following page

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Several state regulations governing the sales or use of e-cigarettes and related products were associated with lower proportions of youth trying or regularly using vaping products, a new study found.

Restricting sales of electronic vapor products to minors, however, was not linked to a lower risk of vaping among teens. "It may be too soon to tell if the state level restrictions are having an impact," said lead author Sarah A. Keim, PhD, of Nationwide Children’s Hospital in Columbus, Ohio, in an interview. "However, it was reassuring to see these early indicators that they may be having an effect so early on, and so these findings were not surprising."

Dr. Keim and her associates investigated possible associations between various state laws related to vaping products, all passed before 2015, and youth use of the products. They relied on 2015 data from 35 state-specific surveys of youth regarding use of vaping products and from the Youth Risk Behavior Survey from the Centers for Disease Control and Prevention, a nationally representative, biannual survey of students in grades 9-12. The Tobacco Control Laws Database of the American Nonsmokers’ Rights Foundation provided information on state laws related to electronic vapor products.

Among the 200,513 teens whose responses were included in the study, 44% had ever used any kind of electronic vapor product. Rates were similar between girls and boys for ever having tried one or currently using one, Dr. Keim reported at the Pediatric Academic Societies annual meeting.

Experimentation began young for most: 35% of respondents tried an e-cigarette before age 14 years, and 18% under age 14 currently use vaping products. By age 17, half of all kids had tried an e-cigarette or related product, and a quarter were currently using them.

The researchers looked at associations with each of the following types of laws:
• Statewide prohibition of vaping products on school property or in workplaces, which includes Arizona, New Hampshire, Vermont, and Virginia for schools and North Dakota for workplaces.
• Prohibition of sales to minors under age 18 years, present in 24 states.
• Prohibition or restriction of sales of e-cigarette products from vending machines, present in 17 states.
• Prohibition or restriction of self-service displays of vaping products, present in 11 states.
• Prohibition or restriction of sampling of electronic vapor products, present in Arizona, Delaware, Kentucky, Maryland, New Hampshire, North Carolina, Oklahoma, and South Carolina.

For most of the regulations, teens had a reduced likelihood of trying or currently using vaping products after adjusting for age, ethnicity, grade level, race, region, and sex. Risk of ever trying a vaping product was 12% lower in states that prohibited their use on school grounds or in workplaces, 6% lower in states that barred sales to those under age 18, and 7% lower in states that restricted or prohibited self-service vaping displays.

The risk of youth currently using electronic vapor products was 5% lower in states with the school grounds and workplace restrictions, and 13% lower in states that restricted self-service displays. Laws restricting minor sales were unrelated to the risk of current vaping among youth. Restricting vending machine sales of vaping products had no association with the risk of a teen ever trying vaping, but it was linked to a 7% lower risk of current use of the products among teens. All these associations were statistically significant based on confidence interval values.

A statistically significant risk increase in vaping use occurred for teens in states that restricted or outlawed sampling of vaping products.

Patients with functional GI disorders had a lower mean ACT score, compared with those without (112 vs. 15; \( P = .03 \)). Functional GI disorders also were associated with higher anxiety scores (34 vs. 14; \( P < .01 \)). Asthma control significantly predicted the presence of functional GI disorders in univariate analysis (odds ratio, 0.9; 95% confidence interval, 0.80-0.99; \( P = .03 \)). However, this significance was lost in a multivariate analysis that adjusted for asthma control, anxiety, and sex. The multivariate analysis revealed continued significant associations between functional GI disorders and anxiety (OR, 1.1; 95% CI, 1.01-1.10; \( P < .01 \)) and female sex (OR, 3.3; 95% CI, 1.00-10.56; \( P < .05 \)).

Dr. Colman speculated that the apparent association of asthma with chronic GI distress could reflect asthma-related inflammation that exacerbates the GI disorders.

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EHR price alert doesn’t reduce lab orders

BY M. ALEXANDER OTTO
Frontline Medical News

Displaying Medicare allowable fees in the electronic health record at the time of order entry did not significantly reduce the number of inpatient lab tests at three Philadelphia hospitals.

In a study involving 98,529 patients and 142,921 admissions, Medicare payment information popped up randomly in the EHR when standard tests including complete blood cell counts, metabolic panels, and liver function tests were ordered. The costs of the labs varied depending on their extent. The message mentioned that “the dollar amount represents Medicare reimbursement for the test. Actual costs to the consumer may vary by patient insurance status.” Just over a third of the patients were actually on Medicare; most had private insurance.

The idea of the study was to see if cost information would curb unnecessary testing, and save money. “There is growing interest in using price transparency to influence medical decision making toward higher-value care,” Mina Sedrak, MD, and her colleagues said in a paper presented at the annual meeting of the Society of General Internal Medicine. It didn’t work out that way. Four tests ordered per patient-day when the messages appeared, and 2.34 when they did not. With messaging, the mean lab fee per patient-day was $38.85, versus $27.59 without it. In an adjusted analyses comparing the intervention to the control group, there were no significant changes in overall test ordering (0.05 tests ordered per patient-day, \( P = .06 \)) or associated fees when pricing information was displayed ($0.24 per patient-day, \( P = .47 \)).

In a subset analysis, the investigators did find a small decrease in orders for the most expensive labs and a small but significant increase in orders for the least expensive ones when physicians aware of cost (top quartile of tests based on fee value: -0.01; \( P = .04 \); bottom quartile: 0.03, \( P = .04 \)).

Despite the overall negative results, there’s still a likely role for cost information in value improvement programs; what the study shows is that...

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VIEW ON THE NEWS

Michael E. Nelson, MD, FCCP, comments: One also needs to consider the effects of information overload and alert fatigue, both of which have been well-documented since the advent of EMRs. Most interesting is the fact that knowledge of the price actually was associated with a slight increase in test ordering, although not statistically significant. It would be even more interesting to conduct a similar study providing the knowledge to both the patient and the physician.
there’s a better way to use it, according to Dr. Sedrak, currently of the City of Hope Comprehensive Cancer Center in Duarte, Calif., and colleagues.

The investigators made several suggestions when reviewing their work. “First, the price transparency intervention in this study was always displayed regardless of the clinical scenario. The presence of this information for appropriate tests may have diminished its impact when tests were inappropriate. Future efforts may consider more selective targeting of price transparency.” It might also be a good idea to price out different testing options for providers, and use actual charges and other more on-point forms of cost estimates, they said, instead of Medicare fees that have little to do with what many patients are actually charged. Targeting only the most expensive tests might also help (JAMA Intern Med. 2017 Apr 21. doi: 10.1001/jamainternmed.2017.1144).

The investigators also noticed a problem when labs are ordered to repeat automatically; clinicians did not see the price information every day, and so missed cost information “when it would be most salient.”

The mean age in the study was 54.7 years; 52% of the patients were white, 39% black, and 57% women. The mean length of stay was about 6 days, and over 80% of the patients were discharged home.

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Proposal would exempt most from MACRA/QPP

BY GREGORY TWACHTMAN

More than half of physicians could be spared from participating in Medicare’s new value-based payment programs in 2018, thanks to a Centers for Medicare & Medicaid Services proposal exempting some physicians.

The proposed 2018 update to the Quality Payment Program (QPP), the payment system created as part of the Medicare Access and CHIP Reauthorization Act (MACRA), would increase the low-volume threshold for participation, exempting practices that receive $90,000 or less in Medicare Part B payments or have 200 or fewer Medicare patients. These would be exempt from participation in either the Merit-based Incentive Payment System (MIPS) or Advanced Alternative Payment Model (APM) tracks of the QPP.

According to the proposed rule, released June 20, the CMS estimates that approximately 572,000 eligible clinicians would be required to participate in MIPS in the 2018 MIPS performance period. After restricting the population of eligible clinician types who are not newly enrolled, the proposed increase in the low-volume threshold is expected to exclude 585,560 clinicians who do not exceed the low-volume threshold.

The CMS is estimating there will be 554,846 MIPS-eligible clinicians in payment year 2020, and most of them will receive either a positive or neutral payment adjustment because of their participation.

Overall, 96.6% of MIPS-eligible physicians will engage in quality reporting in 2020, with 96.1% receiving either a bonus to their Medicare Part B payments or no adjustment, according to CMS estimates. For all eligible clinicians, 76.8% will receive a bonus payment, with all payment bonuses totaling $673.3 million, while those losing money will see their Medicare payments reduced by $173.3 million. The overall aggregate impact will be a 0.9% increase in Part B payments to clinicians.

However, different practice sizes will have different experiences. For example, practices with 1-15 eligible clinicians (114,424 total eligible clinicians in this group) will see in the aggregate a 0.7% increase, while practices with 16-24 eligible clinicians (22,296) will see a 0.4% increase in the aggregate. Practices of 100 or more clinicians (318,841) stand to see the biggest bump in their Medicare payments, with a 1.4% bonus based on the provisions in the proposal.

Ten percent of practices with 1-15 MIPS-eligible clinicians and 10.9% of practices with 16-24 MIPS-eligible clinicians are estimated to receive a decrease in their Medicare payments based on the proposal, while 0.8% of clinicians in practices of 100 or more are expected to see the penalty.

Comments on the proposed update to the QPP are due to the CMS by Aug. 21, 2017.

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Dear CHEST Leaders, Members, and Friends:

The Forum of International Respiratory Societies (FIRS) is an organization comprised of the world’s leading international professional respiratory societies presenting a unifying voice to improve lung health globally. Its members are: the American College of Chest Physicians (CHEST), American Thoracic Society (ATS), Asian Pacific Society of Respirology (APSR), Asociación Latino Americana De Tórax (ALAT), European Respiratory Society (ERS), International Union Against Tuberculosis and Lung Diseases (The Union), the Pan African Thoracic Society (PATS), the Global Initiative for Chronic Obstructive Lung Disease (GOLD), and the Global Initiative for Asthma (GINA). FIRS has more than 70,000 professional members; the physicians and patients they serve magnify our efforts, allowing FIRS to speak for lung health on a global scale.

FIRS is working with the World Health Organization and the United Nations to make sure lung health is represented in national health agendas. FIRS’ position paper on electronic nicotine delivery systems was presented at a side-event at the United Nations High-Level Meeting (UNHL) in New York in 2014 and is now a world standard. At the recent World Health Assembly meeting (May 2017) in Geneva, FIRS launched its Global Impact of Lung Disease report that called for a global clean air standard, strong anti-tobacco laws, and better health care for patients with respiratory disease. FIRS will be reviewing the new WHO Global Air Quality Guidelines and will help promote them globally through advocacy and messaging, as well as by providing air quality expertise. FIRS will be involved at the Coimbra meeting (Sept 26-29) on improving the urban environment, the Monte-video UN High-Level (UNHL) meeting on chronic disease (Oct 18-20), and the UN Ministerial Meeting in Moscow on tuberculosis, and it is preparing for the 2018 UNHL meetings on antibiotic drug resistance, tuberculosis, and chronic diseases.

At the World Health Assembly, FIRS proclaimed September 25 as World Lung Day and hopes to use this as a rallying point for advocacy related to respiratory health or air quality. Lung Disease is the only major chronic disease that does not have a World Day. FIRS produced a Charter for Lung Health (www.firsnet.org/publications/charter) and hopes to have 100,000 persons sign on to it. FIRS also seeks to have lung-health organizations sign on and develop activities that can be carried out to celebrate lung health. Uruguay was the first country to sign the charter. The logos of the organizations who have signed the charter are on the FIRS website at firsnet.org. Activities being planned include editorials, newsletters, and letters-to-the-editor articles, legislative proclamations, social media exposure, and free spirometry, smoking cessation guidance, and carbon monoxide testing, but FIRS is looking for many more ways to celebrate healthy lungs on September 25 and many more partners!

Sixty-five million people suffer from chronic obstructive pulmonary disease and 3 million die of it each year, making it the third leading cause of death worldwide; 10 million people develop tuberculosis and 1.4 million die of it each year, making it the most common deadly infectious disease; 1.6 million people die of lung cancer each year, making it the most deadly cancer; 334 million people suffer from asthma, making it the most common chronic disease of childhood; pneumonia kills millions of people each year, making it a leading cause of death in the very young and very old. At least 2 billion people are exposed to toxic indoor smoke; 1 billion inhale polluted outdoor air; and 1 billion are exposed to tobacco smoke, and the tragedy is that many conditions are getting worse. We cannot sit still and allow this to happen.

FIRS proposes a multipronged campaign to combat lung disease to bring together all people concerned with lung health. It starts with naming September 25 World Lung Day and calling on respiratory health organizations to pledge to improve lung health and help identify ways to celebrate this day.

Please sign up, and share this call for action with your professional, advocacy, and social networks, and those of your friends and families. Please do your part as global citizens to improve lung health. To do so, organizations should indicate they wish to sign on and send their logo to Betty Sax, FIRS Secretariat, betty.sax@ersnet.org. Organizations should also encourage individuals to sign on and show that they are committed to increasing awareness and action to promote global lung health.

Thank you.

Gerard Silvestri, MD, MS, FCCP
CHEST President
Darcy Marciniuk, MD, FCCP
CHEST FIRS Liaison
CHEST Membership News

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Point accrual started on July 5, so you've already been earning points. If you are an FCCP, you began with 30 points awarded for becoming FCCP—that's only 20 points away from the first tier of prizes. To accrue or redeem points, you must be an active member and current with your dues.

Log in to your CHEST account, and access Participation Points in the left column to see your points.

Start earning more points today! Learn more at chestnet.org/participationpoints.

Vaccination: An Important Step in Protecting Health

Patients with chronic lung conditions, like COPD and asthma, need to take extra steps to manage their condition and ensure the healthiest possible future. One important step that may not always be top of mind is vaccination, which can protect against common preventable diseases that may be very serious for those with respiratory conditions. CDC recommends adults with COPD, asthma, and other lung diseases get an annual flu vaccine, as well as stay up to date with pneumococcal and other recommended vaccines. Additional vaccines may be indicated based on age, job, travel locations, and lifestyle.

COPD and asthma cause airways to swell and become blocked with mucus, making it hard to breathe. Certain vaccine-preventable diseases can make this even worse. Adults with COPD and asthma are at increased risk of complications from influenza, including pneumonia and hospitalization. They are also at higher risk for invasive pneumococcal disease and more likely to develop infections including bacteremia and meningitis.

Each year, thousands of adults needlessly suffer, are hospitalized, and even die of diseases that could be prevented by vaccines. Despite increased risks, less than half of adults under 65 years with COPD and asthma have received influenza and pneumococcal vaccination (National Health Information Survey 2015).

Find the latest recommended adult immunization schedule at www.cdc.gov/vaccines/hcp/adults.
CRITICAL CARE COMMENTARY

Conscience Rights, Medical Training, and Critical Care

BY ANA-MARIA DUMITRU, PHD; BENJAMIN W. FRUSH, MA; CHRIS RADLICZ, MS, MPH; PHILIP ALLEN, BS; MARTIN T. BROWN, BS; JEREMY BANNON, BSC; AND JOHN Y. RHEE, MPH

“’No provision in our Constitution ought to be dearer to man than that which protects the rights of conscience against the enterprises of the civil authority.’” – Thomas Jefferson


What is the proper role of conscience in medicine? A recent article in the New England Journal of Medicine (Stahl & Emmanuel. N Engl J Med. 2017; 376(14):1380) is the latest to address this question. It is often argued that physicians who cite conscience in refusing to perform requested procedures or treatments necessarily infringe upon patients’ rights. However, we feel that these concerns stem from a fundamental misunderstanding of what conscience is, why it ought to be respected as an indispensable part of medical judgment (Genuis & Lipp. Int J Family Med. 2013; Epub 2013 Dec 12), and how conscience is oriented toward the end goal of health, which we pursue in medicine.

By failing to define “conscience,”

Continued on following page

EDITOR’S NOTE:

When I invited Dr. Wes Ely – the coauthor of a recent article regarding physician-assisted suicide – to write a Critical Care Commentary on said topic, an interesting thing happened: he declined and suggested that I invite a group of students from medical schools across the country to write the piece instead. The idea was brilliant, and the resulting piece was so insightful that I invite you to read on. Out of that effort, the idea for the present piece was born. The result is an opportunity to hear the students’ voices, not only to stimulate discussion on conscientious objection in medicine but also to remind the ICU community that our learners have their own opinions and that through dialogues such as this, we might all learn from one another.

Lee Morrow, MD, FCCP

 NEWS FROM CHEST

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Long-acting beta,2-adrenergic agonists (LABAs) increase the risk of asthma-related death. Data from a large placebo-controlled US study that compared the safety of another LABA (salmeterol) or placebo added to usual asthma therapy showed an increase in asthma-related deaths in patients receiving salmeterol. This finding with salmeterol is considered a class effect of all LABAs, including indacaterol, one of the active ingredients in UTIBRON NEOHALER.

The safety and efficacy of UTIBRON NEOHALER in patients with asthma have not been established. UTIBRON NEOHALER is not indicated for the treatment of asthma.

Please see additional Important Safety Information, including BOXED WARNING, and Brief Summary of Prescribing Information on adjacent pages.

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Conscience ought to be respected as an indispensable part of medical judgment.

NEWS FROM CHEST
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respiratory tract infection, pneumonia, diarrhea, headache, gastroesophageal reflux disease, hyperglycemia, rinitis. Postmarketing Experience: The following additional adverse reactions of angioedema and dysphonia have been identified during worldwide post-approval use of indacaterol/glycopyrrolate at doses higher than the recommended dose. Because this reaction is reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate the frequency or establish a causal relationship to drug exposure.

DRUG INTERACTIONS: Adrenergic Drugs: If additional adrenergic drugs are to be administered by any route, they should be used with caution because the sympathetic effects of the two drugs may be additive. A component of UTIBRON NEOHALER, non-Potassium-Sparing Diuretics: The electrocardiographic (EKG) changes produced by diuretics as a result of the administration of non-potassium-sparing diuretics (such as loop or thiazide diuretics) can be accentuated when used with beta-agonists. Indacaterol, a component of UTIBRON NEOHALER, is especially when the recommended dose of the beta-agonist is exceeded. Although the clinical relevance of these effects is not known, caution is advised in the coadministration of UTIBRON NEOHALER with non-potassium-sparing diuretics. Monoamine Oxidase Inhibitors, Tricyclic Antidepressants, QTc-Prolonging Drugs: Indacaterol, one of the components of UTIBRON NEOHALER, can interact with other beta-agonists, should be administered with extreme caution to patients being treated with monoamine oxidase inhibitors, tricyclic antidepressants, or other agents known to prolong the QTc interval because the action of adrenergic drugs on the cardiovascular system may be poteniated by these agents. Drugs that are known to prolong the QTc interval may have an increased risk of ventricular arrhythmias.

Beta-Blockers: Beta-adrenergic receptor antagonists (beta-blockers) and UTIBRON NEOHALER may interact with the effect of each other when administered concurrently. Beta-blockers not only block the therapeutic effects of beta-agonists, but may produce severe bronchospasm in COPD patients. Therefore, patients with COPD should not normally be treated with beta-blockers. However, under certain circumstances, e.g., as prophylaxis after myocardial infarction, there may be no acceptable alternatives to the use of beta-blockers in patients with COPD. In this setting, caution is advised and beta-blockers should be considered, although they should be administered with caution.

Anticholinergics: There is potential for an additive interaction with concomitantly used anticholinergic medications. Therefore, coadministration of UTIBRON NEOHALER with other anticholinergic-containing drugs may be considered to increase in anticholinergic adverse effects. Inhibitors of Cytochrome P450 3A4 and P-g Efflux Transporter: Drug interaction studies with indacaterol, a component of UTIBRON NEOHALER, were carried out using potent and specific inhibitors of CYP3A4 and P-g (i.e., ketoconazole, erythromycin, verapamil, and rifampin). The data suggest that systemic clearance of indacaterol is influenced by modulation of both P-g and efflux transporters. In the 2-fold area under the curve (AUC) increase caused by the strong dual inhibitor ketoconazole reflects the impact of maximal combined inhibition. Indacaterol was evaluated in clinical trials for up to 1 year in doses up to 600 mcg. Monitoring of the key contributors of indacaterol clearance, CYP3A4 and P-g, has no impact on safety of therapeutic doses of indacaterol. Therefore, no dose adjustment is warranted at the recommended dose of UTIBRON NEOHALER when administered concurrently with inhibitors of CYP3A4 and P-g. USE IN SPECIFIC POPULATIONS: Pregnancy: Teratogenic Effects: Pregnancy Category C. There are no adequate and well-controlled studies in pregnant women. Animal reproduction studies were conducted with individual components, indacaterol and glycopyrrolate. Because animal reproduction studies are not always predictive of human response, UTIBRON NEOHALER should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Women should be advised to contact their physician if they become pregnant while taking UTIBRON NEOHALER. Indacaterol: Indacaterol was not teratogenic in Wistar rats and New Zealand rabbits at approximately 340 and 770 times, respectively, the MRHD in adults (an AUC basis at maternal subcutaneous doses up to 1 mg/kg/day in rats and rabbits). Glycopyrrolate: Glycopyrrolate was not teratogenic in Wistar rats or New Zealand White rabbits at approximately 1400 and 350 times, respectively, the MRHD in adults (an AUC basis at maternal inhaled doses up to 3.83 mcg/ml per day in rats and up to 4.4 mcg/kg/day in rabbits). Non-teratogenic Effects: Indacaterol: There were no effects on perinatal and postnatal development in rats at approximately 110 times the MRHD in adults (in an AUC basis at maternal subcutaneous doses up to 0.3 mg/kg/day). Glycopyrrolate: There were no effects on perinatal and postnatal development in rats at approximately 1100 times the MRHD in adults (in an AUC basis at maternal subcutaneous doses up to 1.88 mcg/kg/day). Labor and Delivery: There was no evidence of adverse events in any of the two human trials that have investigated the effects of UTIBRON NEOHALER during labor and delivery. Because beta-agonists may potentially interfere with uterine contractility, UTIBRON NEOHALER should be used during labor only if the potential benefit justifies the potential risk. In human parturients undergoing Cesarean section, 96 minutes after a single intramuscular injection of 0.166 mcg/kg glycopyrrolate, umbilical plasma concentrations were low. Nursing Mothers: UTIBRON NEOHALER: It is not known whether UTIBRON NEOHALER is excreted in human breast milk. Because many drugs are excreted in human milk, caution should be exercised when UTIBRON NEOHALER is administered to a nursing woman. Since there are no data from well-controlled human studies on the use of UTIBRON NEOHALER by nursing mothers, based on the data for the individual components, a decision should be made whether to discontinue nursing or to discontinue UTIBRON NEOHALER, taking into account the importance of UTIBRON NEOHALER to the mother. Indacaterol: It is not known whether indacaterol is excreted in human breast milk. Glycopyrrolate (including its metabolites) have been detected in the milk of lactating rats and reached up to 10-fold higher concentrations in the milk than in the blood of the dam. Pediatric Use: UTIBRON NEOHALER is not indicated for use in children. The safety and efficacy of UTIBRON NEOHALER in pediatric patients have not been established. Geriatric Use: Based on available data, no adjustment of UTIBRON NEOHALER dosage in geriatric patients is warranted. UTIBRON NEOHALER can be used at the recommended dose in elderly patients 72 years of age and older. Of the total number of subjects in clinical studies of UTIBRON NEOHALER, 45% were aged 65 and older; 11% were aged 75 and older. No overall differences in safety or effectiveness were observed between these subjects and younger subjects. Further clinical studies of UTIBRON NEOHALER in patients with severe renal impairment estimated GFR less than 30 mL/min/1.73 m2 or end-stage dialysis disease requiring dialysis, UTIBRON NEOHALER should be used if the expected benefit outweighs the potential risk since the systemic exposure to glycopyrrolate may be increased in this population. Hepatic Impairment: Based on the pharmacokinetic characteristics of its monotherapy components, UTIBRON NEOHALER can be used at the recommended dose in patients with mild to moderate hepatic impairment. Studies in subjects with severe hepatic impairment have not been performed. OVERDOSAGE: In COPD patients, doses of up to 600/124.8 mcg UTIBRON NEOHALER were inhaled over 2 weeks and there were no relevant effects on heart rate, QT interval, ventricular tachycardia, or any other cardiac adverse effects. There were no overt signs of toxicity. In patients with a single subcutaneous dose of 1 mg/kg for up to 24 hours, the maximum reported symptom was dizziness reported in 3 of 17 patients. In a study with Xanodyne Derivatives, Steroids, or Diuretics: Treatment of overdose consists of discontinuation of UTIBRON NEOHALER together with institution of appropriate supportive measures. Indacaterol, one of the components of UTIBRON NEOHALER, is a beta-agonist and a beta-receptor blocker may be considered, bearing in mind that such medicine can produce bronchospasm. Cardiac monitoring is recommended in cases of overdosage. Indacaterol is a beta-agonist and with overdose of indacaterol are those of excessive beta-adrenergic stimulation and occurrence or exaggeration of any of the signs and symptoms, e.g., angina, hypotension, including marked tachycardia, up to 200 bpm, arrhythmias, nervousness, headache, tremor, dry mouth, palpitation, muscle cramps, nausea, vomiting, diarrhea, dizziness, fatigue, malaise, hypo/hyperpyrexia, metabolic acidosis and insomnia. As with all inhaled sympathomimetic medications, cardiac arrest and even death may be associated with an overdose of indacaterol. In COPD patients, single doses of indacaterol 3000 mcg were associated with moderate increases in pulse rate, systolic blood pressure and QTc interval. Glycopyrrolate: An overdose of glycopyrrolate may lead to anticholinergic signs and symptoms such as nausea, vomiting, dizziness, lightheadedness, blurred vision, increased intraocular pressure (causing pain, vision disturbances or reddening of the eye), obstipation or difficulties in voiding. In COPD patients, repeated orally inhaled administration of glycopyrrolate at total doses of 124.8 mcg and 249.6 mcg once-daily for 28 days were well tolerated. PATIENT COUNSELING INFORMATION: Advise the patient to read the FDA-approved patient labeling (Medication Guide and Instructions for Use).

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NEWS FROM CHEST

The actions we perform – and those we omit – constantly shape our individual consciences.

Theor Med Bioeth

Oxford, Oxford University Press.

That's the place of conscience altogether.

By attributing appropriate value to an individual's conscience, we thereby recognize the centrality of conscience to identity and personal integrity. Consequently, we see that forcing an individual to impinge on his/her conscience through coercive

meants significantly violates that person's autonomy and dignity as a human being capable of moral decision-making.

In the practice of medicine, the free exercise of conscience is especially relevant. When patients and physicians meet to act in the pursuit of the patient's health, they begin the process of conscience-mediated shared decision-making, rife with the potential for disagreement. Throughout this process, a physician should not violate a patient's conscience rights by demanding a procedure or treatment that the physician cannot perform in good conscience. Moreover, to insert an external arbiter (e.g., a professional society) to resolve the situation by means of contradiction of conscience would have the same violating effect on one or both parties. One common debate is to the application of conscience in the setting of critical care focuses on the issue of physician-assisted suicide and euthanasia (PAS/E) (Rhee J, et al. Chest. 2017;152[3]3]. Accepted for Sept 2017 publication). Those who would deny physicians the right to conscientiously object to PAS/E depict this as merely an issue of the physician's personal preference. Given the distinction between pre-
erence and conscience, however, we recognize that much more is at play. For students and practitioners who hold that health signifies the “well-working of the organism as a whole,” (Kass L. Public Interest. 1975; 40(summer):11-42) and feel that the killing of a patient is an action that goes directly against the health of the patient, the obligation to participate in PAS/E represents not only a violation of our decision-making dignity, but also subverts the critical component of clinical judgment inherent to our profession. The conscientiously practicing doctor who follows what they believe to be their professional obligations, acting in accordance with the health of the patient, may reasonably conclude that PAS/E directly contradicts their obligations to pursue the best health interests of the patient. As such, their refusal to participate can hardly be deemed a simple personal preference, as the refusal is both reasoned and reasonable. Indeed, experts have concluded that regardless of the legality of PAS/E, physicians must be allowed to conscientiously object to participate (Goligher et al. Crit Care Med. 2017; 45(2):149).

As medical students who have recently gone through the arduous medical school application process, we are particularly concerned with the claim that if one sees fit to exercise conscientious objection as a practitioner, they should leave medicine, or choose a field in medicine with few ethical dilemmas. To crassly exclude students from the pursuit of medicine on the basis of the shape of their conscience would be to unjustly discriminate by assigning different values to genuinely held beliefs. A direct consequence of this exclusion would be to decrease the diversity of thought, which is central to medical innovation and medical progress. History has taught us that the frontiers of medical advancement are most ardently pursued by those who think deeply and then dare to act creatively, seeking to bring to fruition what others deemed impossible. Without conscience rights, physicians are not free to think for themselves. We find it hard to believe that many physicians would feel comfortable jettisoning conscience in all instances where it may go against the wishes of their patients or the consensus opinion of the profession.

Furthermore, as medical students, we are acutely aware of the importance of conscientious objection due to the extant hierarchical nature of medical training. Evaluations are often performed by residents and physicians in places of authority, so students will readily subjugate everything from bodily needs to conscience in order to appease their attending physicians. Evidence indicates that medical students will even fail to object when they recognize medical errors performed by their superiors (Madigosky WS, et al. Acad Med. 2006; 81(1):94).

It is, therefore, crucial to the proper formation of medical students that our exercise of conscience be safeguarded during our training. A student who is free to exercise conscience is a student who is learning to think independently, as well as to shoulder the responsibility that comes as a consequence of free choices. Ultimately, we must ask ourselves: how is the role of the physician altered if we choose to minimize the role of conscience in medicine? And
do patients truly want physicians who forfeit their consciences even in matters of life and death? If we take the demands of those who dismiss conscience to their end – that only those willing to put their conscience aside should enter medicine – we would be left with practitioners whose group think training would stifle discussion between physicians and patients, and whose role would be reduced to simply acquiescing to any and all demands of the patient, even to their own detriment. Such a group of people, in our view, would fail to be physicians.

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July 4th was bittersweet for me, this year. Independence days of my childhood were spent grilling, sitting by the campfire on the lakes and rivers of Northern Michigan, watching the fireworks turn the night sky red, white, and blue. These fond memories were a painful reminder that others like me may not have the privilege to experience such joy, secondary to their background.

I don’t remember the first time that I heard the tale of my parents coming to America. They were both medical students from India, who received brightly colored brochures from American hospitals inviting them to come further their medical training. Due to the deficit of physicians in the United States, the hospitals even loaned money to medical students, so they would do their residencies in America. My parents took advantage of this opportunity and embarked on a journey that would define their lives. Often, my mother would talk about my father leaving for the hospital on Friday morning only to return to his wife and two toddlers on Monday afternoon. As a child, I remember my uncles taking bottles of milk to the hospital to make chai to fuel through their grueling overnight calls. These immigrant tales were the backdrop of my childhood, the basis of my understanding of America. I was raised in an immigrant community of physicians who were grateful for the opportunities that America offered them. They worked hard, reaped significant rewards, and substantially contributed to their communities. Maybe, I am just nostalgic for my childhood, but this experience, I believe, is still an integral part of the American dream.

The recent choice to restrict immigration from specific nations is disturbing at best and reminiscent of an America that I have never known. More than 7,000 physicians from Libya, Iran, Somalia, Sudan, Syria, and Yemen are currently working in the United States, providing care for more than 14 million people. An estimated 94% of American communities have at least one doctor from one of the targeted countries. These physicians are more likely to work in rural and underserved communities and provide essential services. They are immigrants who have come to America to better their lives and, in turn, have bettered the lives of those around them. They are my parents. Not all physicians are good people or are worthy of the American dream, but America is a better place for welcoming those who are willing to work hard to make a better life for themselves. An important criticism of the effect of migration of medical professionals to the United States is that it has a positive impact on the communities they work in.
States has been the loss of human capital to their respective nations, but never the ill-effect they have had on the nations they have emigrated to.

The 2015 Educational Commission for Foreign Medical Graduates (ECFMG) reported that a quarter of practicing physicians in the United States are international medical graduates (IMGs) and a fifth of all residency applicants were IMGs. Measuring the impact of the IMGs who have come to America is difficult to quantify but can be assessed by countless anecdotes and success stories. Forty-two percent of residency applicants were IMGs. Graduates (IMGs) and a fifth of all physicians in the United States are foreign born. Twenty-eight American Nobel prize winners in Medicine since 1960 are immigrants and taking their parents had accomplished. The American College of Chest Physicians (CHEST), over the past 15 years, has had several Presidents who are American immigrants. One of them, Dr. Kalpalatha K. Guntpalli, President 2009-2010, I have met, and I was humbled by the experience. She is brilliant, kind, and modest and without her knowing, she has served as one of the role models for my career.

I applaud CHEST for standing with other member organizations to oppose the immigration hiatus (Letter to John F. Kelly, Secretary of Homeland Security, Feb 7, 2017). The medical organizations made four concrete proposals:

- Reinstate the Visa Interview Waiver Program, as the suspension of this program increases the risk for significant delays in new and renewal visa processing for trainees from any foreign country;
- Remove entry restrictions of physicians and medical students from the seven designated countries that have been approved for J-1, H-1B or F-1 visas;
- Allow affected physicians to obtain travel visas to visit the United States for medical conferences, as well as other medical and research related events; and
- Prioritize the admission of refugees with urgent medical needs who had already been checked and approved for entry prior to the executive order. These recommendations were good but not broad enough. The decision to bar immigration for any period of time, from any country, is an affront to the American dream with long-lasting consequences, most importantly, the loss of health-care services to the American populace. My Congressman knows how I feel about this, does yours?

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FIGURE 1
Foreign-Born Share of Healthcare Workers by Occupation, 2010

Source: Adapted from American Community Survey 5 year estimates (2010-2014) and IPUMS-USA, University of Minnesota, www.ipums.org.

<table>
<thead>
<tr>
<th>Occupation</th>
<th>Foreign-Born Share</th>
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<tbody>
<tr>
<td>Physicians &amp; Surgeons</td>
<td>27.8%</td>
</tr>
<tr>
<td>Nursing, Psychiatric &amp; Home Health Aides</td>
<td>21.9%</td>
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<tr>
<td>Registered Nurses</td>
<td>14.7%</td>
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<tr>
<td>Therapists</td>
<td>12.2%</td>
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<tr>
<td>Therapists</td>
<td>12.2%</td>
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<tr>
<td>Healthcare Support</td>
<td>12.7%</td>
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Catching Up With Our CHEST Past Presidents

Where are they now? What have they been up to? CHEST’s Past Presidents each forged the way for the many successes of the American College of Chest Physicians, leading to enhanced patient care around the globe. Their outstanding leadership and vision are evidenced today in many of CHEST’s strategic initiatives. Let's check in with Dr. Goldberg.

Allen I. Goldberg, MD, Master FCCP
President 1998-1999

I arrived in Toronto in 1998 to start my term as President of the American College of Chest Physicians. (I had always loved Toronto, where I had spent months training in pediatric critical care at “Sick Kids” [Toronto’s Children’s Hospital] and collaborating with Audrey King on disability issues and public policy in Ontario.) CHEST 1998 in Toronto was equally exciting. What I remember - with humility – was that being CHEST President is not about “you.” It is about “The President,” who is honored and revered by all members for what CHEST truly represents … excellence in healthcare education, communication, and information. Everyone came up to me to respect and honor the role … including awesome Past Presidents who lovingly shared their insights and experience and others (including many who became future presidents) to volunteer their assistance. I was in awe of these leaders and how they demonstrated selfless service.

And so I began my year of presidential service leadership. What I remember best is the respect all around the world for CHEST and what it does to unite people into actions that improve health globally. The President serves CHEST members to facilitate working together, which makes a difference. My presidential year culminated in the 65th anniversary conference in Chicago in 1999. All year, I had worked with my mentor (C. Everett Koop, MD, FCCP(Hon), to plan an opening ceremony that would be inspirational and unforgettable. For years, we had shared personal/private conversations. This time, we planned to communicate in public to inspire others and help them understand key issues.

In 2000, I began my second term as President. My presidency culminated in the 65th anniversary conference in Chicago in 2001. All year, I had worked with my mentor (C. Everett Koop, MD, FCCP(Hon), to plan an opening ceremony that would be inspirational and unforgettable. For years, we had shared personal/private conversations. This time, we planned to communicate in public to inspire others and help them understand key issues.

In 2002, I retired to work more closely with Dr. Koop and we considered critical for the future of healthcare and global health. Soon after my Presidential term, I took 2 years off for sabbatical to work more closely with Dr. Koop (2000-2002). Then, I retired to continue to focus on our work together and as personal caregiver for my wife, Evi Faure, MD, FCCP. Dr. Koop and I met many times and held more public presentations, including the 2003 Surgeons’ General National Meeting on Overcoming Health Disparities at Howard University arranged with CHEST Past President Dr. Alvin Thomas.

All our joint efforts focused on the importance of Communication in Health Care. We shared the belief that communication of health information would create the “informed patient and family” who would then work together in partnership with health-care professional team members. We thought that this would be the best way to improve and reform health-care delivery. We sought to provide information (the “what”)

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http://www.chestnet.org/News/Blogs/CHEST-Thought-Leaders/2013/06/Dr-Koops-Legacy--Reflections-on-Mentorship
CHEST Joint Congress in Basel, Switzerland

Members of CHEST leadership, faculty, and staff traveled to Basel, Switzerland, in June, to participate in the CHEST Joint Congress, which was co-hosted with the Swiss Respiratory Society, Schweizerische Gesellschaft Fur Pneumologie (SPG). Overall, there were approximately 1,100 total attendees, representing over 40 countries, who enjoyed the scientific program and gained valuable chest medicine knowledge. Among the many topics presented were diagnosis and treatment of ILD; biologics for severe asthma; EBUS for molecular analysis; and ICS in COPD. Plus, hands-on, interactive workshops were offered for learning or reviewing more procedural skills. We invite you to view webcasts of five of the Basel sessions at bit.ly/chestsgp2017.

The CHEST Joint Congress in Basel represented the second collaborative scientific conference endeavor with a third party, the first being the CHEST Conference held in Amsterdam May 6-9, COPD: Current Excellence and Future Development.

New Tools in Campaign to Fight Asthma

The Allergy & Asthma NetWork, the nation’s leading patient education and advocacy organization for people with allergy and asthma, has once again joined forces with the CHEST Foundation in an effort to empower patients suffering from severe asthma.

The campaign’s focus is to educate health-care providers, patients, parents of asthmatics, and the public about the most current treatment options for asthma, highlight the importance of referring to specialists to improve patient outcomes, and bring to light the role of the entire health-care team in the care of a patient with severe or difficult-to-control asthma.

This is the second year of this growing campaign, and there are several new and exciting materials.

Severity Assessment Tool

Available online and in print, the severity assessment tool was designed to help a patient, and the clinician, understand the severity of their asthma. Not only does the tool evaluate the severity of their condition, but it also helps the patient become more aware of their symptoms. The seven-question assessment includes questions on usage of quick-relief or rescue inhalers, visits to the ED/hospital, physical activity, controller medication, and quality of sleep.

Patient and Caregiver Testimonials

The campaign features several patient and caregiver testimonials that tell the stories of patients and caregivers who experienced changes in their health.

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Pulmonary/Critical Care with Sleep Cambridge Health Alliance • Cambridge, MA

Cambridge Health Alliance (CHA) an award-winning public healthcare system, has an opportunity for a Pulmonary/ Critical Care Physician to join our existing Pulmonary team. Our system is comprised of three hospital campuses and an integrated network of both primary and specialty care practices in the Boston area. CHA is a teaching affiliate of both Harvard Medical School (HMS) and Tufts University School of Medicine.

Candidate will practice Pulmonary/CC medicine and ideally incorporate dedicated Sleep Medicine time, as well as possess a strong interest in resident and medical student teaching. Incoming physician should possess excellent clinical/communication skills and a strong commitment to serve our multicultural safety net patient population. This position has both inpatient and outpatient responsibilities. We offer a supportive and collegial environment with a strong infrastructure, inclusive of an electronic medical record system (EPIC). Candidates will have the opportunity to work in a team environment with dedicated colleagues similarly committed to providing high quality healthcare. Our employees receive competitive salary and excellent benefits.

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Health-care weaponization, PTSD, depression in caregivers

**Disaster Response**

**The tragic weaponization of health care**

The Syrian conflict has highlighted the dangers to health-care workers (HCWs) in humanitarian crises. The Lancet-American University of Beirut Commission on Syria reports on the weaponization of health care in Syria—a strategy of depriving people of their health-care needs. Targeting of HCWs was recognized early in the Syrian war with targeting of health-care facilities being frequently reported throughout the conflict. HCWs facing extreme supply shortages have been reported to resort to desperate measures: using urine bags with added anticoagulants for blood collection and crafting homemade external fixators for fractures. Sadly, the Syrian conflict is not unique. The International Committee of the Red Cross (ICRC) documented 2,398 episodes of violence directed at health facilities in 11 countries affected by armed conflict between 2012 and 2014 alone. In Syria and elsewhere, the exodus of trained medical personnel, due to lack of medical training in trauma, emergency medicine, and intensive care, puts populations at further risk in these regions. The International Red Cross and Red Crescent Movement has started the Health Care in Danger (http://healthcareindanger.org) initiative to highlight this weaponization of health, supporting efforts by HCWs to advocate for their rights and their patients’ rights at a global level. This highlights the needs for CHEST members responding to humanitarian crises to ensure they have appropriate training to work in these environments and deploy with an organization that can provide adequate safeguards.

Dr. Maves is a military service member. The opinions expressed herein are his own and do not necessarily reflect the official opinions of the Department of the Navy, Department of Defense, or the US Government.

**Practice Operations**

**The House AHCA /Senate BCRA compared with ACA (Affordable Care Act)**

Health-care costs are a fundamental driver of insurance costs, which leads to challenges to coverage affordability for millions of families. There is ongoing debate whether the current law (Affordable Care Act [ACA/Obamacare]) and the republicans alternatives (American Healthcare Act [AHCA] and Better Care Reconciliation Act [BCRA]) do enough to address the cost challenges. Here is a brief summary of the key similarities and differences.

**Similarities:**
1. Children will be covered up to age 26.
2. Coverage of pre-existing conditions continues (high risk pools will be subsidized by a state government but premiums are up to twice as much as individual coverage).
3. Tax credit (based on age and family size rather than income level).
4. Insurance can charge older customers more than younger (up to 3X under ACA, 5X under AHCA/BCRA).
5. No annual or lifetime payout limit (but states may apply waivers allowing insurers to apply limits).

**Differences:**
1. Insurance will no longer be mandatory (no individual or employer mandates, but there is a 30% increase in premiums for 1 year for not maintaining individual continuous coverage).
2. Medicaid expansion (expanded under ACA to 135% of poverty level income) will stop in 2020.
3. Restriction on “Abortion Funding” (any facility that offers abortion will not receive federal funding) for 1 year.
4. Taxes on health care will be removed (including taxes on prescription drugs, OTC, premiums, and medical devices).
5. Allowing policies for major illness or injury (with elimination of the requirement to cover ten essential health benefits, allowing states to modify).
6. Health-care reform undoubtedly is complicated, and there are a lot of questions in the air about the future of health care under the Trump Administration. Few certainties: change is coming. MACRA is here to stay.

Adel Bassily-Marcus, MD, FCCP

**Networks**

**Posttraumatic Stress Disorder Post-Lung Transplant**

The majority of transplant physicians are mainly concerned with issues posttransplant that are focused on the graft function. But recently, neurocognition and posttransplant psychiatric stress disorder have been found to have significant impact on quality of life and mortality after transplantation. Posttraumatic stress disorder (PTSD) is described as re-experiencing a traumatic event in addition to having avoidant and hyperarousal symptoms, which last for a period of at least 1 month. Studies of PTSD in solid organ transplant recipients have revealed a significantly higher prevalence of PTSD symptoms (10% to 17%) compared with the general population (prevalence of 3.5% to 6%). In one study of heart transplant recipients, patients who met the criteria for PTSD in the first year posttransplant had a higher risk for 3-year mortality (OR=13.74) [Dew et al. J Heart Lung Transplant. 1999;18(6):549-562].

Lung transplant recipients are at a high risk for developing PTSD due to exposure to several traumatic events, such as a life-threatening exacerbation of the underlying lung disease, undergoing transplant surgery, intensive care unit stay, delirium and episodes of infection, and acute and chronic rejection. However, data regarding the prevalence and risk factors for PTSD post-lung transplant are limited.

The prevalence of PTSD post lung transplantation has been reported to

**Thank you to Our Supporters**

The CHEST Foundation and Allergy and Asthma Network would like to thank our generous supporters, AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, and Novartis for making this campaign possible. It is through supporters, who are active participants in helping grow this campaign, that these important materials are able to have an impact on patient outcomes and create long-lasting social change.

To view the campaign materials, visit us at asthma.chestnet.org.

Immunology (ACAAI), the Allergy & Asthma Network, and CHEST Foundation have partnered to develop a shared decision-making tool for adults with severe asthma. This tool will be launched at CHEST 2017 in October. Available online and in print, it was created for patients and clinicians to work together to improve self-management skills, choose the best treatment plan for the patient, and increase adherence. This patient-centered approach in clinical settings improves patient satisfaction of care and overall outcomes.

**Fellow-in-Training Member**

Ryan Maves, MD, FCCP

**Steering Committee Member**

Dr. Maves is a military service member. The opinions expressed herein are his own and do not necessarily reflect the official opinions of the Department of the Navy, Department of Defense, or the US Government.

Benjamin Buckley who was 7 years old at the time of Ben’s passing, he was on a preventive med. He was going to the doctor routinely. We had actually just been to the asthma doctor. We were seeing somebody, had an action plan, and everybody knew what they had to do. Even with all of that, it still came to this. Benjamin still lost his life, and we never knew this was something that could happen,” stated Cristin Buckley, mother of Benjamin Buckley who was 7 years old at the time of his death. These testimonial videos will be used to raise awareness of the condition, and the importance of managing and monitoring symptoms.

**Shared Decision Making Tool**

The American College of Allergy, Asthma, and
be 12.6% to 15.8%. In lung transplant recipients with clinically significant PTSD symptomatology; the presence of symptoms of re-experiencing (29.5%) and arousal (33.8%) were more common than avoiding symptoms (18.4%) [Gries et al. J Heart Lung Transplant. 2013;32(5):525-532]. In another study by Dew et al, in 178 lung transplant recipients, all PTSD occurred in the early months posttransplant with a median duration of symptoms of 12 months (IQR 7.2 to 18.5 months) [Dew et al. Gen. Hosp Psychiatry. 2012;34:127-138]. A higher burden of PTSD is noted in patients who are younger, have a lower income, have a previous history of a traumatic event, and have bronchiolitis obliterans (Gries et al. J Heart Lung Transplant. 2013;32(5):525-532).

The challenges that remain include determining the true prevalence of PTSD in the lung transplant recipient in the LAS era using standard diagnostic criteria, documenting the adverse effects of PTSD on medical compliance, morbidity, and mortality; and developing interventions to mitigate the adverse effects of PTSD through well-designed multicenter prospective studies.

Vivek Ahya, MD
Steering Committee Member

Women’s Health
Caregiver Burden in the ICU and Beyond

Family members of patients in the ICU who transition to the role of caregivers following discharge are at high risk for psychosocial distress. Post-intensive care syndrome-family (PICS-F) describes the symptoms of depression, posttraumatic stress, and anxiety commonly found in this population (Davidson et al. Crit Care Med. 2012;42(2):618-624). Women are more commonly called upon to adopt the role of caregiver for family members with chronic medical conditions or mental illnesses. Worldwide estimates indicate that 57% to 81% of all caregivers are women (Sharma et al. World J Psych. 2016;6(2):7-17).

Family burden begins during the acute phase of critical illness. As surrogate decision-makers, they frequently face decisional conflict and decisional regret, especially in scenarios that limit life-sustaining therapies (Long et al. Curr Opin Crit Care. 2016;22:613-620). The prevalence of PICS-F is high as family members attempt to balance their role in the ICU with personal obligations (Choi et al. J Korean Acad Nurs. 2016;46(2):159-167). Those who perceive that they are not receiving complete information from the medical team, and who do not find their physician comforting, have been shown to suffer a greater symptom burden (Davidson et al).

With the growing older adult population, and increased ICU survival, family members are often called upon to serve as caretakers to the chronically critically ill (Choi et al.). These caregivers have more depressive symptoms, worse health outcomes, and significant professional and personal lifestyle disruptions (Cameron et al. N Engl J Med. 2016;374(19):1831-1841). In many caregivers, depressive symptoms persist at 1 year after ICU admission, with rates comparable to caretakers of patients with dementia (Haines et al. Crit Care Med. 2015;43(5):1112-1120). Caregivers who are younger, female, minorities, and those with pre-existing depression are at especially high risk for worse mental health outcomes (Davidson et al; Cameron et al).

Caregivers of ICU survivors are vulnerable and undersupported. Interventions such as ICU diaries, telephone-based mindfulness exercises, and stress management strategies have shown promise in alleviating PICS-F symptoms (Choi et al.). During the acute ICU stay, how medical providers communicate, and how we help family members make sense of what has happened and their new roles as caregivers have an impact (Davidson et al.). From an individual in a study of psychosocial morbidity in caregivers of ICU survivors: “Leaving the hospital is not the end for some people. The next place is just as hard, sometimes worse” (Haines et al.

Further studies are needed to identify interventions that will truly address this population’s unique needs.

Margaret Pisani, MD, FCCP
Steering Committee Member
Nicole Bournival, MD
Fellow-in-Training Member

2017
Education Calendar

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- **Critical Care Ultrasound: Integration Into Clinical Practice**
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