Scientists from AFIP's Department of Cellular Pathology, Division of Molecular Pathology, have determined the nature of the virus responsible for the "Spanish flu" outbreak that swept the world in 1918-19, killing at least 21 million people worldwide, including over 43,000 U.S. servicemen. It was the worst infectious disease episode ever. Experts have long theorized that the virus was a dangerous "swine" flu, but it wasn't until the AFIP team performed genetic analysis on preserved tissue samples from victims of the flu that the theory could be confirmed. Their findings now "open the door" to discovering why this particular flu was so deadly (Science. 1997;275:1793.)

Flu viruses have been around for centuries, and experts who study them generally agree that they originate in birds and are occasionally transmitted to humans. The result is typically an influenza pandemic. The viruses also change rapidly, and to combat them, new vaccines must be made each year. In 1918, however, influenza viruses hadn't been discovered and vaccines weren't available. That spring it was observed that pigs and pig farmers in Iowa got sick simultaneously. By the fall, a new and deadly flu virus took hold. Victims typically died within a week, their lungs filled with fluid. "No one has ever seen that before or since," says Jeffrey K. Taubenberger, MD, PhD, chief, Division of Molecular Pathology, Department of Cellular Pathology, the lead author of the study. "It was a unique pathology."

Millions died worldwide, including approximately 675,000 Americans. "It killed so many young people that the average life expectancy in the U.S. was reduced by more than 10 years," Taubenberger notes. Among them were thousands of U.S. servicemen mobilized for World War I. Some of the 43,000 who died were autopsied, and their preserved tissue samples were sent to the Army Medical Museum (now the AFIP) for future study.

After the pandemic, came the questions: Why was this flu so devastating? Why did so many die? Theories grew that perhaps the flu had gone directly from birds to pigs, where it mutated into a particularly deadly human virus. By the 1930s, influenza viruses had been discovered, but only an indirect analysis of the serum of 1918 survivors pointed to a "classic swine flu" as the cause.

Over the years, experts have looked at the preserved tissue samples to characterize the virus, but without success. Recent genetic advances, however, led Dr. Taubenberger, molecular biologist Ann Reid, and their colleagues to try again. They began by examining the records of 70 U.S. servicemen who died of the flu. "Since the virus actively replicates for only a few days, we needed tissue samples from a victim who died within 1 week of the onset of symptoms in order to find the Spanish Flu, continued on page 10
New Technologies for the 21st Century

Our most recent successes in discovering clues to the 1918 Spanish Flu reflect the scope of AFIP's unique and changing capabilities.

Seventy-nine years and about 2,000,000 accessioned cases came between that day in 1918 when a young soldier's lung tissue was accessioned and the time the "Spanish Flu" viral RNA sequence was published. During that time, archivists meticulously catalogued tissue, preserved it, and filed it so that when the time came it could be retrieved and studied. Tissue can be located in minutes; today a pathologist can interpret that well-preserved and maintained slide as easily as in 1918.

While assuring that our specimens do not change with time, our staff, and the techniques they use, must change. AFIP's Department of Cellular Pathology staff—both the authors of the Science paper and others, developed and refined methods for investigating archival tissues using molecular techniques that are second to none. A few years ago, the idea of sequencing RNA from 80-year-old paraffin-embedded blocks would have been considered preposterous. Now these, and similar DNA-based techniques, are used not only in a wide variety of research areas—such as breast cancer, soft tissue tumors, hematologic and infectious diseases, but also almost 10,000 times per year in molecular diagnostic assays that are routine at the AFIP.

Creating and maintaining a staff capable of anticipating and meeting the needs of tomorrow is a challenge. AFIP staff includes internationally renowned diagnostic pathologists, microbiologists, molecular biologists, chemists, computer scientists, and engineers. They carry out basic investigations, using techniques ranging from light microscopy to magnetic resonance microscopy and from histochemical staining to laser Raman spectroscopy and develop new methods, in partnership with "traditional" diagnostic pathologists, that improve both our diagnostic abilities and our understanding of disease pathogenesis.

We are confident that our efforts to preserve the old—both our invaluable specimen collection and our diagnostic expertise—and develop new technologies and abilities, will enable AFIP to achieve its vision both now and in the 21st century.

Michael J. Dickerson
Col, USAF, MC
The Director
The AFIP’s course, “Controversias y Adelantos en Patologia Quirurgica,” presented entirely in Spanish, will be held at the Parque Balneario Hotel, Santos, Sao Paulo, Brazil, August 20-24, 1997. Joining AFIP course directors Cesar A. Moran, Maj, USAF, MC; Clara S. Heffess, COL, MC, USA; and Florabel G. Mullick, MD, SES, Director of AFIP’s Center for Advanced Pathology, will be Marcello Franco, MD, director of surgical pathology at the University of Sao Paulo and president of the Brazilian Society of Pathologists. Dr. Franco will serve as associate director of the course.

“This course has been offered three previous times—one in Puerto Rico, with 176 attendees; once in Valencia, Spain, with 166 attendees; and most recently in Guadalajara, Mexico, with a record 257 in attendance,” says Dr. Mullick. “We anticipate another huge turnout in Santos. The course has clearly demonstrated that Latin American and Spanish pathologists want to strengthen their ties to the AFIP, and we look forward to another successful program in August.”

Microscopes will be available at no cost to the participants, who are encouraged to bring their difficult cases for a second opinion from the course faculty. Joining noted experts from the AFIP on the faculty will be representatives from the University of Texas - Southwestern Medical School, Dallas, Texas; Wayne State University, Detroit, Michigan; Baylor College of Medicine, Houston, Texas; M.D. Anderson Cancer Center, Houston, Texas; University of Granada, Spain; and Mount Sinai Medical Center, Miami, Florida.

New to the 1997 course will be an overview of forensic pathology. Other topics set for discussion include: endocrine pathology, otolaryngic pathology, genitourinary pathology, gynecologic and breast pathology, dermatopathology, pulmonary pathology, soft tissue pathology, neuropathology, gastrointestinal pathology, and cytology.

Santos, located in the state of Sao Paulo, is Brazil’s largest port city. It is a popular tourist destination in August, with a number of attractive beaches and a pleasant climate.

For more information about the course, contact Mr. Carlos Moran at (202) 782-2556, by e-mail at Moran@email.afip.osd.mil, or by FAX at (202) 782-7166.

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Customer Satisfaction Surveys mailed to contributors

Ten thousand post cards containing an important customer satisfaction survey will be mailed with consultation reports starting in April and continuing until the supply is exhausted. Contributors are asked to complete the survey and return it postage-paid to AFIP. The survey will provide AFIP with valuable information on how our customers feel about our service. This survey will be compared to ones conducted in 1992 and 1994 to see if our customers feel we have made an improvement in our consultation service. The survey is also available on the AFIP web site (WWW.AFIP.MIL). For more information, contact Mr. Frank Roberts, Quality Assurance Coordinator, at (202) 782-2695, or by e-mail at ROBERTSF@email.afip.osd.mil.
William Inskeep II, COL, VC, USA
appointed Chair, Department of Veterinary Pathology

William A. Gardner, Jr, MD, elected Chair, Scientific Advisory Board
Elson B. Helwig MD, receives Distinguished Pathologist Award

Upon his arrival at the Armed Forces Institute of Pathology in 1946, Dr. Helwig was appointed chair, Department of Skin and Gastrointestinal Pathology, and registrar of the American Registry of Dermal Pathology, serving from 1947 to 1980. He also served as chair, Center for Advanced Pathology, and associate director for consultations from 1955 until 1977.

During this period, Dr. Helwig pioneered the field of dermatopathology as a key founder of the specialty and offered a very popular, thorough, and highly coveted program in dermatopathology known as the Osborne Fellowship. In 1980, he turned to his full-time professional activities as chair, Department of Gastrointestinal Pathology.

During his 44 years at the AFIP, Dr. Helwig has trained many pathologists and dermatopathologists, given hundreds of lectures and seminars nationally and internationally, and authored or coauthored over 200 scientific reports and 7 books. He has served as visiting professor at many universities and holds the rank of clinical professor at the Uniformed Services University of the Health Sciences.

Dr. Helwig has actively served as a member of the editorial boards of several journals and as a member of committees, boards, and panels of national and international organizations, including the American Academy of Dermatology, American Society of Dermatopathology, American Joint Committee for Cancer Staging and End Results, American Board of Pathology, National Research Council, National Academy of Sciences, International Academy of Pathology, and the World Health Organization.

He has received numerous distinguished honors, awards, and medals, including the Presidential Distinguished Federal Civilian Service Medal, Distinguished Civilian Service Awards from the Department of the Army and Department of Defense; and the first John Shaw Billings Lifetime Achievement Award from the AFIP. One of his most cherished awards was the honorary degree of Doctor of Sciences presented by the Indiana University at the dedication ceremonies for the university’s new medical research and library building. This was in recognition of his lifetime of distinguished service to the academic and medical science communities.
PROFILE

Sarah S. Frankel, MD, named winner of 1997 Castleman Award

Dr. Frankel is a graduate of Wellesley College and Rush University Medical College, Chicago. She completed an internship in general surgery and a year of residency training in otorhinolaryngology at the Albert Einstein College of Medicine in New York, and a residency in anatomic pathology at the New York Hospital-Cornell Medical Center in 1990.

Previous assignments included staff positions at The New York Hospital and The Manhattan Eye and Throat Hospital, and as assistant professor of pathology at Cornell University Medical College. In 1991, she moved to Buffalo, New York, where she was on the staff of the Buffalo General Hospital and the Roswell Park Cancer Center. She also served as assistant professor of pathology and ophthalmology at the State University of New York at Buffalo, where she twice received the Siegal Commendation for Excellence in Teaching.

In 1994, Dr. Frankel assumed her present position as staff pathologist at AFIP. She is also a clinical assistant professor of pathology at Georgetown University School of Medicine and serves as a staff scientist in the Division of Retrovirology, Department of Vaccine Development, Walter Reed Army Institute of Research.

REPOSITORY AND RESEARCH SERVICES

FAXes replace case acknowledgement post cards

Due to the success of our initiative to fax case acknowledgement forms to contributors at the time a case is accessioned and the Institute’s efforts in decreasing case turnaround times, the Receiving and Accessions Division will no longer be mailing contributors a case acknowledgment post card. The faxing of case acknowledgment forms has substantially improved communications with contributors by quickly providing them the case accession number once assigned, as well as the name and phone number of the department that will be reporting on the case, information not available on the post card. If this change will cause a problem for any of our contributors, we ask that they contact the administrator, Repository and Research Services, at (202)782-2500.

The Receiving and Accessions Division will also no longer routinely mail x-rays back to contributors, unless they request them at the time of case accessioning. This is the same policy currently in effect for the return of paraffin blocks. This new initiative will not only save postage but will eliminate confusion. The x-rays not immediately requiring return after the case has been finalized will remain on file at the Institute and will be returned upon written/faxed request to our Information Release Office. Anyone having questions concerning this change can contact the above number.
Museum archivist keeps history alive

Rhode answers many questions from the public about the history of medicine. "I'd say we get about three calls or letters a day, sometimes more. People also come in to do research. We ask that they call or write a few weeks in advance, so we can be sure we can help them."

He also occasionally writes exhibits. "I wrote the Civil War medical illustration cases that are on exhibit now. This is actually the fourth version of it that I've written. We have to keep changing the photographs and watercolors, so they won't fade. Hopefully, after it closes, we'll be able to get this exhibit on the Museum's web site, since the artwork won't be displayed again for years," says Rhode.

Does he have a favorite piece? "If I do have one at the moment, it certainly changes over time. I wouldn't pick one of our best-known pieces like 'The Deathbed of Abraham Lincoln' sketch by Museum artist Hermann Faber. I've seen that too many times. I like to find something new. I'm partial to the photographic collections, though," he replied.

Rhode has recently been working on a Civil War medical photography book. "Dr. Blair Rogers, a New York City plastic surgeon, suggested we work on one together years ago. I've written chapters on the Museum and our photographers. Hopefully we'll wrap it up and get it to Duke University for publication this fall. Meanwhile, I've written an introduction to Photographic Atlas of Civil War Injuries, a reprint of many of the Museum's photographs."

Redding and Rhode are also working on a Guide to the Collections for the Museum, expected to be published by the American Registry of Pathology. They are hoping to select enough interesting and exciting illustrations to make it attractive to individuals and not just to libraries.

Recently, Rhode has been lecturing on the Museum and its photographic work. "In the past six months, I guess I've spoken four or five times on this. I did a short version for the Washington Society for the History of Medicine in the fall and a longer version for the Smithsonian in February. Fortunately, I can use all the research I did for the book in my talks."

For more information or to use the Otis Archives, contact Rhode at (202) 782-2212, or by e-mail at Rhode@email.afip.osd.mil.

Private Julius Fabry was wounded in August 1864. His leg was amputated at the knee. When his femur became infected, Dr. George Otis, the Museum's second curator, performed the difficult operation of amputating Fabry’s leg at the hip. Photographed by Museum staff photographer, E. J. Ward. SP 276.
Portable DNA analysis system delivered to AFIP

By Elizabeth Campos Rajs
Lawrence Livermore Laboratory

A portable DNA analysis system that will allow military specialists to quickly identify human remains in the field, test food and water in remote locations for contamination, and identify pathogenic bacteria on the battlefield was recently delivered to the AFIP by Lawrence Livermore Laboratory.

"Developed by a team of Lab scientists in the Micro Technology Center, the DNA analysis system is revolutionary because of its small size," said M. Allen Northrup, a biomedical engineer and the principal investigator. The hand-held unit operates on 1.5 watts — compared to the laboratory-sized version that consumes 1,000 watts — and the whole system fits in a small suitcase.

"This has wide-ranging applications to the military and society in general," said Victor W. Weedn, LTC(P), MC, USA, Chief Deputy Medical Examiner, Office of the Armed Forces Medical Examiner, and Program Manager, Department of Defense DNA Registry.

Weedn assisted the Defense Advanced Research Project Agency in overseeing this project during the last 3 years.

As a chief deputy in the U.S. government's only medical examiner's office, Weedn has been called around the world to identify human remains. He also was called on to help identify the remains of Czar Nicholas in Russia.

"This is the beginning of a revolution in instrumentation and DNA testing," said Victor W. Weedn, LTC(P), MC, USA, Chief Deputy Medical Examiner, Office of the Armed Forces Medical Examiner, and Program Manager, Department of Defense DNA Registry.

"This is the time in the world that there's been a product produced for portable DNA testing," he said. "This instrument permits the most rapid, sensitive method of DNA testing available. It permits specific identification of a target molecule as well as quantification of the target molecules, and this instrument could be applied to any DNA testing."

Performing a technique known as polymerase chain reaction (PCR), the machine makes millions to billions of copies of specific DNA from traces of blood or other cells — whether plant, animal, or germ — at a fraction of the time.

It also performs real-time quantitative detection of the DNA as it is being synthesized, a feature unique to this instrument.

"In experiments in collaboration with Roche Molecular Systems in Alameda and the University of Maryland Medical School, the miniature instrument was used to detect HIV and Hepatitis C virus in human patient samples in less than 20 minutes," Northrup said. "Extremely low viral load levels were detected which has important implications for monitoring therapy of such diseases."

In the past, DNA samples taken in the field had to be sent back to the laboratory for analysis. Now, analysis and identification can take place anywhere in the world in a matter of minutes.

"The portable unit can do it more consistently with more specificity and in a fraction of the time," Weedn said. "I'm thrilled to be a part of this."

For the last 3 years, Northrup's project has been funded by the Department of Defense's Advanced Research Projects Agency's Microelectromechanical Systems Program, under the direction of Ken Gabriel. Turning the unit over to the Army concludes that project, although the team will continue to improve and test the technology with continued DOD funding.

"Without this hardworking critical team, the project wouldn't have been successful. A lot of credit goes to them," Northrup said.

"The portable DNA analyzer has broad applications," Northrup explained. "It can be used for clinical diagnosis of infectious diseases and cancers, identifying pathogenic bacteria in the environment, and testing agricultural products for diseases. It has potential for broad application for almost all biological analyses and research."

Weedn plans to display one of the machines in the National Museum of Health and Medicine of the AFIP.

"I truly believe in my heart of hearts that this PCR-microchip technology is revolutionary," he said.

Rapid DNA testing may become more common thanks to this portable DNA analysis system.
AFIP on Global Environmental Collaboration: International Program on Environmental Health and Toxicologic Studies of Arsenosis, Fluorosis, and Thallium Endemic Areas in Southwest Guizhou Province, People’s Republic of China

Recently, Jose A. Centeno, PhD, a research scientist from the Department of Environmental and Toxicologic Pathology, and four other American scientists were invited to visit Guizhou Province in China to study health problems associated with arsenic, fluoride, and thallium exposures originating from geological materials, particularly high-arsenic coal. The U.S. Task Force was formed as a multidisciplinary, multiagency effort with representatives from the Armed Forces Institute of Pathology, the U.S. Geological Survey, the U.S. Environmental Protection Agency, the University of Arizona, and the Rocky Mountain Poison and Drug Center.

“Health problems associated with high levels of toxic trace elements and metals, including arsenic, fluoride, mercury, and thallium, are known to afflict millions of people in several countries,” Centeno points out. “Endemic arsenosis cases have been recently reported in India, Bangladesh, Taiwan, Japan, and Inner Mongolia. In the United States, exposures to arsenic have been reported in several states, including Nevada, Michigan, and Utah.”

The influence of geologic materials in human health is an area of fertile scientific and medical research requiring the development of international collaborations in order to understand the distribution, mobilization, and potentially hazardous effects of these materials. Since the early 1950s, more than 3,000 cases of endemic arsenicism induced by burning of high-arsenic coals (As > 1,000 mg/kg) in Southwest Guizhou Province, P.R. China, have been reported. In this region, the coal is locally mined and burned indoors in an open fire without a chimney. The incidence of arsenicism, fluorosis, selenosis, and thallium poisoning cases have been correlated with high levels of these toxic elements and metals found in these coals.

The U.S. delegation was invited by the National Nature Science Foundation of China, the Academy of Sciences, and the Institute of Geochemistry of China. In collaboration with the Chinese institutions and provincial authorities, Dr. Centeno and his colleagues conducted on-site studies on the extent of the environmental problem and on the assessment of clinical features associated with the presence of arsenic, fluoride, and thallium, and established a dialogue for the development of a memorandum of agreement to conduct medical and toxicologic studies. The environmental health assessment program includes chemical characterization studies, toxicology and clinical treatment, and the collection of clinical and pathological specimens to study the effect(s) of these contaminants in tissue.

“Results from this project could provide useful information to U.S. military medicine by improving our understanding of the human health impact of these contaminants and their distribution, mobility, and behavior in environmental geochemical sites where American troops could be potentially deployed,” Centeno says.

The presence of arsenic in these coals in the range of 10 to perhaps 10,000 ppm provides an excellent opportunity to contribute to knowledge of the dose response of arsenic. This is a major issue being addressed by the Environmental Protection Agency for the upcoming reauthorization of the Clean Water Act, and has particular relevance to the elevated levels of arsenic in drinking water in eastern Michigan, India, and Bangladesh.
Spanish Flu, Continued from page 1
evidence we needed,” Taubenberger notes.
Most did not meet the criteria, but a
few did, and one case in particular led the
team to believe the virus was still replicat­ing
when the victim died. Sections from
that case revealed focal acute bronchiolitis
and alveolitis with individual cell necrosis
of bronchiolar epithelial cells, indicative
of viral pneumonia.
He was a 21-year-old private who
died at Fort Jackson, South Carolina 5
days after being infected. “He was
completely healthy with no medical
history until he got the flu,” Taubenberger
says. He and Ann Reid then utilized
polymerase chain reaction (PCR) tech­ni­ques to amplify minute amounts of the
victim’s genetic material for sequence
analysis.
Nine fragments of viral RNA were
sequenced from the coding regions of five
influenza genes: hemagglutinin,
neuraminidase, nucleoprotein, matrix 1,
and matrix 2. The sequencing was
particularly difficult to accomplish since
over the years the viral RNA had broken
down into minute pieces only 200 nucleo­tides long. The technique to isolate RNA
from formalin-fixed, paraffin-embedded
tissues was developed in the AFIP’s
Division of Molecular Pathology, and the
RNA from this case was extracted by
research biologists Amy E. Krafft, PhD,
and Karen E. Bijwaard.
Ann Reid devised methods to amplify
small fragments of the 1918 viral genes
using degenerate PCR primers that would
recognize conserved sequences in human,
swine, and avian influenza viruses. Once
sequences were obtained, she was able to
devise PCR primers that more closely
matched the sequences from the 1918 flu.
“The hemagglutinin gene matches
closest to swine influenza viruses,
showing the first direct evidence that the
virus came into humans from pigs,”
Taubenberger says. Phylogenetic analyses
of the gene fragments from the 1918 virus
were performed by research biologist
Thomas G. Fanning, PhD, using sophisti­cated computer software packages that
compare differences in related genes.
What does the future hold?
Taubenberger estimates that his team has
analyzed about 7% of the virus, but he’s
confident that continued studies will
provide more information about the flu’s
complete genetic code. “On two other
occasions since 1918, a major
flu outbreak has been clearly
caused by the appearance of
new influenza strains that
mutated in pigs
—the 1957
Asian flu and
the 1969 Hong
Kong flu,” he
points out, “and
in 1976 there
was an isolated
outbreak of a
new swine flu
virus in
humans.
Millions of dollars were spent to vaccine
the population out of fear this could have
been as damaging as the 1918 flu. While
that particular strain turned out not to be
lethal, the 1918 results confirm that ‘swine
flus’ remain a significant public health
threat. Understanding the genetic make-up
of this flu may help us combat future
epidemics and prevent such a terrible loss
of life.”

Tri-Service School of Histotechnology

The Tri-Service School of Histotechnology, (Department of Scientific Laborato­ries), is located on the grounds of Walter Reed Army Medical Center at the Armed
Forces Institute of Pathology, Washington, D.C. The school has a staff consisting of
a program director, course superintendent, dean of civilian students, two
instructors, and eight adjunct faculty members who teach up to 25 students per
course. Students are trained in preparing tissue specimens for macroscopic and
microscopic examination, postmortem procedures, microtomy, embedding, routine
hematoxylin and eosin (H&E) staining and special stains to further pathologic
diagnosis.
The school is undergoing major changes to its curriculum to enhance the
quality of academia and training of the students to become proficient and quali­fied histotechnicians. Initially, the Tri-Service School offered two courses on a
yearly basis, but the new curriculum will consist of 3 months of academic study
and 6 months of actual hands-on training under the supervision of the school’s
instructors and adjunct faculty members. The school trains military students from
the Air Force, Navy, and Army along with civilians either directly out of high
school or individuals wishing to change their perspective career fields.
Currently, the school is undergoing approval for accreditation by the National
Accrediting Agency for Clinical Laboratory Sciences (NAACLS), which allows
students to test for certification through the American Society of Clinical Patholo­gist (ASCP) to be certified as Histotechnician (HT) ASCP. For additional
information about the Tri-Service School of Histotechnology, please forward
correspondences to AFIP, Bldg. 54, Rm 2107, 6825 16th Street, NW, Washington,
D.C. 20306-6000, call (202) 782-2801, or FAX: (202) 782-7663.
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**ABSTRACTS OF RECENT PUBLICATIONS BY AFIP STAFF**

**Carcinoids of the jejunum and ileum: an immunohistochemical and clinicopathologic study of 167 cases**

Allen P. Burke, M.D. Rebecca M. Thomas, M.D. Al M. Elsayed, M.D. Leslie H. Sobin, M.D.

**BACKGROUND.** Carcinoid tumors of the gastrointestinal tract differ in their clinical and histopathologic features, depending on the site of origin. There are few clinicopathologic studies that specifically describe jejunoileal carcinoid tumors.

**METHODS.** One hundred sixty-seven ileal and jejunal carcinoids were retrospectively studied with emphasis on clinical, pathologic, immunohistochemical, and prognostic features.

**RESULTS.** The mean age of patients at the time of presentation was 62 + 12 years (range, 13-93 years). Eight patients had carcinoid syndrome (5%) and 1 had Zollinger-Ellison syndrome. Twenty-six percent of tumors were multiple, and 77% were transmurally invasive; 31% had regional lymph node metastases only, and 32% had liver or mesenteric metastases. Ninety-three percent of tumors had an insular growth pattern. Serotonin was expressed in 86% of tumors (86 of 102), chromogranin in 92%, and neuron specific enolase in 95%. Twenty percent of tumors (10 of 51) expressed prostatic acid phosphatase; 96% were argentaffin, and 98% argyrophil. Of 80 cases with follow-up data (mean follow-up, 52 + 5 months), 21% were dead of disease, 16% were dead of other causes, 19% were alive with disease, and 44% had no evidence of disease at last follow-up. The 5-year Kaplan-Meier survival estimate for all cases was 58%. By univariate analysis, survival was negatively correlated with distant metastases at the time of surgery (P = 0.002), mitotic rate (P = 0.01), tumor multiplicity (P = 0.01), the presence of carcinoid syndrome (P = 0.02), depth of invasion (P = 0.03), and female gender (P = 0.05); by multivariate analysis, survival was negatively associated with distant metastasis (P = 0.002), carcinoid syndrome (P = 0.01), and female gender (P = 0.03).

**CONCLUSIONS.** Jejunoileal carcinoid tumors have a relatively high rate of transmural invasion and aggressive clinical behavior. They are usually insular and largely argentaffin, with a high rate of chromogranin and serotonin positivity. These features differentiate jejunoileal carcinoids from other gastrointestinal carcinoids.

**Inflammatory pseudotumor of lymph nodes: a study of 25 cases with emphasis on morphological heterogeneity**

Cesar A. Moran, MD, Saul Suster, MD, and Susan L. Abbondanzo, MD

The clinicopathological and immunohistochemical findings in 25 cases of inflammatory pseudotumor of lymph nodes (115) are presented. The patients were 13 women and 12 men between 8 and 81 years of age. Clinically, symptoms of prior infection, fatigue, abdominal pain, weight loss, fever of unknown origin, pelvic inflammatory disease, or nausea and night sweats were obtained in 15 patients, whereas six patients presented with asymptomatic lymphadenopathy. In four additional patients, no clinical information was obtained. The involved nodes included cervical, supraclavicular, inguinal, mesenteric, and mediastinal lymph nodes. In two cases, there was synchronous involvement of separate lymph node groups (inguinal and cervical in one case and cervical and mediastinal in another case), whereas in a third patient there was synchronous involvement of the spleen and a paraaortic lymph node. Histologically, the lesions were characterized by a fibrosing/inflammatory process that showed marked heterogeneity and striking variation from case to case. Based on their histological features, the lesions could be classified into three different groups: Stage I was characterized by the appearance of single or multiple small foci containing a spindle cell proliferation admixed with a prominent inflammatory background, with complete preservation of the remainder of the nodal architecture; stage II was characterized by more diffuse involvement of the lymph node with a marked inflammatory response admixed with a prominent myofibroblastic proliferation leading to subfocal effacement of the nodal architecture, often with extension of the process beyond the capsule into perinodal fat; and stage III was characterized by almost complete replacement of the lymph node by diffuse sclerosis with scant residual inflammatory elements and total loss of the normal nodal architecture. Immunohistochemical studies in 20 cases showed a striking number of vimentin- and actin-positive myofibroblastic cells with moderate increase in CD20/CD45* small lymphocytes and polyclonal plasma cells in the stage I lesions, the emergence of numerous CD68+ histiocytes admixed with lymphocytes, plasma cells, and abundant fibromyofibroblastic cells in the stage II lesions, and only few remaining scattered CD68+ histiocytes and fibroblasts in the stage III lesions. Our findings suggest that inflammatory pseudotumor of the lymph node represents an evolving, dynamic process that may adopt different morphological appearances depending on its stage of evolution. Recognition of the various stages of this process may be of importance for differential diagnosis with other fibrosing/inflammatory conditions of lymph nodes.

**Detailed RT-ISPCR protocol for preserving morphology and confining PCR products in routinely processed paraffin sections**

Yan-gao Man, Zhengping Zhuang, Gary L. Brathauer, Omar Bagasra and Fattaneh A. Tavassoli

A modified reverse transcription in situ polymerase chain reaction (RT-ISPCR) protocol has been developed for mRNA detection in routinely processed paraffin sections. Compared to previously published methods, this protocol applies several new strategies, including (I) placing tissue sections on regular microscopic slides to reduce the size of the reaction chamber; (2) substituting enzymatic digestion with overnight incubation at 80°C to eliminate “cross-links” induced by formalin fixation; (3) adding anti-AE1/AE3 antibodies to RT solution to create a sponge-like tissue surface to trap and confine newly formed PCR products; (4) reducing the number of high-temperature treatment steps to avoid complete tissue desiccation; and (5) placing sections at 4°C after RT reaction and amplification to facilitate newly formed PCR products to settle down. With these strategies, we have been able to obtain reproducible results in which the morphology of tissue is well preserved, different cell types are readily distinguishable, and the amplified PCR products are easily identifiable. This protocol can also be used for detection of DNA in paraffin sections with stronger signals and better preservation of morphology. The specificity of this method is confirmed by results from different controls and from other approaches, including immunohistochemistry with specific antibodies, in situ hybridization with specific mRNA probes, and tube PCR with the same primers and PCR conditions used for RT-ISPCR. The basic rationale of our approach is discussed.

**Cell Vis.** 1996;3:389-396.

**Hum Pathol.** 1997;28:332-338.
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