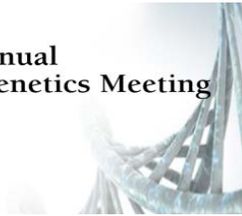




2017 | ACMG Annual
Clinical Genetics Meeting



Developing Care Models for Patients with Secondary Genomic Findings

Held in Phoenix, Arizona, March 21-25, 2017

Date of Release: April 3, 2017

Expiration Date: April 3, 2020 (CME, NSGC, P.A.C.E.®)

Estimate Time of Completion: 2 hours

Course must be completed by the expiration dates

COURSE DESCRIPTION

Routine pursuit of incidental and secondary genomic findings within DNA sequences has prompted a need for the development of better structured clinical management approaches of these results. Screening of DNA sequence in search of these findings constitutes a precursor to an eventual use of genomics for population screening; therefore the WHO criteria for screening that calls for the availability of “facilities for diagnosis and treatment” is relevant to these efforts. The clinical approach to patients ascertained in this manner will initially require extrapolation from patient data ascertained “traditionally”; however sites taking on the management of this new set of patients need to acquire data specific to this ascertainment approach so that management can be refined over time. This session will review four separate efforts to approach patients with secondary genomic findings. The panel will discuss clinical management issues including institutional support for infrastructure, as well as efforts to develop standardized approaches to clinical follow-up care.

Dr. Leslie Biesecker (Delivery of Secondary Findings to ClinSeq Participants at the NIH Clinical Center) declined to release his slides but you will be able to listen to his presentation.

LEARNING OBJECTIVES

At the conclusion of this course, participants should be able to:

- Describe approaches to the management of secondary genomic findings
- Outline knowledge gaps in this emerging area of clinical genomics
- Examine growing data around penetrance and expressivity of secondary genomic findings
- Compare patient care models across institutions and specialties

TARGET AUDIENCE

All healthcare professionals interested in the diagnosis, management, treatment and prevention of genetic conditions and increasing their understanding of the genetic basis of common, chronic health problems affecting both children and adults will find the programming applicable to their practice. These select sessions from the ACMG Annual Meeting are targeted for the following professionals:

- Medical and clinical geneticists
- Physicians of all specialties with an interest in genetics, genomics and the genetic basis of disease
- Genetic counselors
- Laboratory geneticists, directors, technicians and technologists
- Researchers
- Pathologists
- Educators
- Nurses
- Dietitians
- Physician assistants
- Biotechnology and pharmaceutical development professionals
- Fellows, Trainees and Students
- Public health professionals
- Genetic/consumer advocates
- Others with an interest in the science and art of medical genetics and genomics

SESSIONS

- Delivery of Secondary Findings for Multiple Conditions to A Precision Health Cohort at Geisinger.- Michael Murray, MD, FACMG, FACP
- Delivery of Secondary Findings to ClinSeq Participants at the NIH Clinical Center- Leslie Biesecker, MD, FACMG
- Delivery of Secondary Findings Associated with Cancer Risk at University of Pennsylvania- Katherine Nathanson, MD, FACMG
- Delivery of Secondary Findings Associated with Risk for Structural Heart Disease at Johns Hopkins- Cynthia James, ScM, PhD, CGC

Accreditation:

The American College of Medical Genetics and Genomics is accredited by the Accreditation Council for Continuing Medical Education (ACCME) to provide continuing medical education for physicians.

Credit Designation:

The American College of Medical Genetics and Genomics designates this activity for a maximum of 2 *AMA PRA Category 1 Credits*[™]. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

Genetic Counselor Credit

The National Society of Genetic Counselors (NSGC) has authorized American College of Medical Genetics and Genomics to offer up to 2 Category 1 contact hours for this OnDemand course. The American Board of Genetic Counseling (ABGC) will accept CEUs earned for this course for the purposes of genetic counselor certification and recertification. Reporting of credits is sent to NSGC quarterly. Additional fee (~\$25) applies for NSGC credit that is billed by NSGC.

P.A.C.E. CEUs – Laboratory Directors and Laboratory Personnel

ACMG is approved as a provider of continuing education programs in the clinical laboratory sciences by the American Society for Clinical Laboratory Science (ASCLS) Professional Acknowledgment for Continuing Education (P.A.C.E.®) Program. The American College of Medical Genetics and Genomics designates this course for a maximum of 2 contact hours. ACMG is approved by the Florida Board of Clinical Laboratory Personnel as CE Provider. ACMG is approved by the California Department of Health Services through the ASCLS P.A.C.E.® Program as CE Provider #275.

HIPAA Compliance

The ACMG supports medical information privacy. While the ACMG is not a “covered entity” under HIPAA 1996 and therefore is not required to meet these standards, ACMG wishes to take reasonable steps to ensure that the presentation of individually identifiable health information at ACMG-sponsored events has been properly authorized. All presenters have completed a form indicating whether they intend to present any form of individually identifiable healthcare information. If so, they were asked either to attest that a HIPAA-compliant consent form is on file at their institution, or to send ACMG a copy of the ACMG HIPAA compliance form. This information is on record at the ACMG Administrative Office and will be made available on request.

Content Validation

ACMG follows the ACCME policy on Content Validation for CME activities, which requires:

Content Validation and Fair Balance

1. ACMG follows the ACCME policy on Content Validation for CME activities, which requires:
 - a) All recommendations involving clinical medicine must be based on evidence that is accepted within the profession of medicine as adequate justification for their indications and contraindications in the care of patients.
 - b) All scientific research referred to, reported or used in CME in support or justification of patient care recommendations must conform to the generally accepted standards of experimental design, data collection and analysis.
2. Activities that fall outside the definition of CME/CE; “Educational activities that serve to maintain, develop, or increase the knowledge, skills, and professional performance and relationships that a physician uses to provide services for patients, the public, or the profession” (source: ACCME and AMA) will not be certified for credit. CME activities that promote recommendations, treatment, or manners of practicing medicine or pharmacy that are not within the definition of CME/CE or, are known to have risks or dangers that outweigh the benefits or, are known to be ineffective in the treatment of patients.
3. Presentations and CME/CE activity materials must give a balanced view of therapeutic options; use of generic names will contribute to this impartiality. If the CME/CE educational materials or content includes trade names, where available, trade names from several companies must be used.

Off-label Uses of Products

When an off-label use of a product, or an investigational use not yet approved for any purpose, is discussed during an educational activity, the accredited sponsor shall require the speaker to disclose that the product is not labeled for the use under discussion, or that the product is still investigational. Discussions of such uses shall focus on those uses that have been subject of objective investigation.

Disclaimer: *ACMG educational programs are designed primarily as an educational tool for health care providers who wish to increase their understanding of the application of genomic technologies to patient care. The ACMG does not endorse, or recommend the use of this educational program to make patient diagnoses, particular by individuals not*

trained in medical genetics. Adherence to the information provided in these programs does not necessarily ensure a successful diagnostic outcome. The program should not be considered inclusive of all proper procedures and tests or exclusive of other procedures and tests that are reasonably directed at obtaining the same results. In determining the propriety of any specific procedure or test, a healthcare provider should apply his or her own professional judgment to the specific clinical circumstances presented by the individual patient or specimen.

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Members of the ACMG Staff, Education and Program Committees involved in planning the 2017 ACMG Annual Clinical Genetics Meeting are required to disclose relevant relationships which could be perceived by some as a real or apparent conflict of interest in planning. All disclosures have been reviewed and conflicts of interest resolved by the Education Committee COI sub-committee or the Executive Director and CME Associate Director and conflicts of interest are disclosed. In the cases where a conflict existed then the committee member refrained from the discussion.

Following is a list of program and education committee members who have disclosed one or more such relationships and names of companies with which those relationships exist:

EC = Education Committee; PC = Program Committee; S = ACMG Staff

1. Major stockholder/ownership interest	6. Non-remunerative positions of influence such as officer, board member, trustee, or public spokesperson (All Committee Members Below are on ACMG Committees –Members with other affiliations are listed)
2. Grant/Research Support (External)	7. Receipt of intellectual property
3. Salary/Employment/Royalty(ies)/Honoraria	8. Other
4. Consultant/consulting fees/other remuneration	
5. Speakers' bureau	

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SPEAKERS AND MODERATORS

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Moderator/ Speaker: Michael Murray, MD, FACMG, FACP

Director of Clinical Genomics, Geisinger Genomic Medicine Institute

Financial relationships to disclose. (Self): Advisory for Merck, Invitae, Regeneron



Dr. Murray is a clinical geneticist who directs the GenomeFIRST program at Geisinger.

Moderator/ Speaker: Leslie Biesecker, MD, FACMG

Chief and Senior Investigator, National Human Genome Research Institute

Financial relationships to disclose: (Self): Receives honoraria from Genentech, Wiley-Blackwell



Dr. Biesecker is a clinical and molecular geneticist and Chief of the Medical Genomics and Metabolic Branch and Director of the Physician-Scientist Development Program at the National Human Genome Research Institute of the National Institutes of Health, which he joined in 1993. He uses genetic and genomic technologies to study the etiology of genetic disorders and has published nearly 300 primary research articles, reviews, and chapters. He

received his medical training at the Univ. of Illinois, Pediatrics at the Univ. of Wisconsin, and Clinical and Molecular Genetics at the Univ. of Michigan. His laboratory has elucidated the etiology and natural history of numerous diseases. In addition, he developed the ClinSeq® program, which began clinical genomics research in 2006, before the widespread availability of next generation sequencing. He co-directs a CLIA-certified molecular diagnostic laboratory within NHGRI. Dr. Biesecker serves as an editor or board member for several biomedical journals, was a member of the board of directors for the American Society of Human Genetics, is an advisor to the Illumina Corporation, and served on the advisory panels for the World Trade Center and Hurricane Katrina victim identification efforts. He was recently elected to the National Academy of Medicine of the National Academy of Science.

Speaker: Katherine Nathanson, MD, FACMG
Professor, University of Pennsylvania
No financial relationships to disclose.



Katherine (Kate) L. Nathanson, MD is a Professor of Medicine, in the Division of Translational Medicine and Human Genetics at the Perelman School of Medicine of the University of Pennsylvania. She also is Associate Director for Population Sciences and Chief Oncogenomics Physician in Abramson Cancer Center. Dr. Nathanson received her BA from Haverford College and MD from the University of Pennsylvania School of Medicine. She completed residencies in both Internal Medicine and Clinical Genetics, along with a post-doctoral fellowship in cancer genetics. Dr. Nathanson has published over 220 peer reviewed articles in journals such as Nature, JAMA, New England Journal of Medicine and Cancer Cell. Dr. Nathanson has an extensive record of national service, serving on committees for multiple organizations, such as ACMG and AACR, several editorial boards, and scientific review committees including for the National Institutes of Health. She has been elected to the American Society of Clinical Investigation and the American Association of Physicians.

Speaker: Cynthia James, ScM, PhD, CGC

Assistant Professor, Department of Medicine, Division of Cardiology, Johns Hopkins

No financial relationships to disclose.



Cynthia James, ScM, PhD is an Assistant Professor of Medicine at Johns Hopkins University in Baltimore, Maryland. She is a board-certified genetic counselor with a doctorate in Human Genetics. She works in the Center for Inherited Heart Diseases (Division of Cardiology). Her research is focused on 1) psychosocial adaptation to inherited cardiac disease, 2) the role of genetic and environmental factors on disease expression, particularly in patients and families with arrhythmogenic right ventricular cardiomyopathy, and 3) informing genetics / genomics policy through studying the experiences of families.

Participation Instructions

1. Participant logs into ondemand.acmg.net
2. Once logged in the participant will access the session they would like to view. They will be asked if they would like to claim credit for the meeting, or if they will not claim credit for the meeting. Then, this information (Course description) will appear, and participant will have to attest that they have read the information. They will then click Continue.
3. After that, the participant will be able to select the credit types to claim.
4. For each session with a post-test, the participant will need to mark and complete the matching pre-test.
5. Then the participant will watch the session presentations.
6. Participant will complete viewing all session content. "Check marks" indicate which presentations have been viewed.
7. After viewing all presentations within a session, participant will click the "Claim Credit for Session" button under the CME dropdown at the top of the page.
8. Participant should take and then successfully pass the post-test. If they do not pass with a score of 80% or higher, they will have unlimited tries to pass the post-test.
9. Participant will continue the steps above to earn credits for additional sessions.
10. If a session does not have a test attached, the participant will not need to take a pre- or posttest, but will have to complete a Concurrent or Plenary session-specific evaluation to claim credit.
11. To print their certificate, the participant will click the "Print Certificate" button under the CME dropdown at the top of the page. Participant must complete the meeting evaluation (one time only) before they can access their certificate. Participant will then choose their certificate(s). The certificate(s) will be automatically updated as they earn new credits.

Stream Requirements

Network

For best results, use a hardwired network connection instead of wireless

Full Screen Viewing

If you would like to view the webcast full screen, display the tool bar at the bottom and click the double arrow in the far right corner. The screen will enlarge to the full screen of your system. To restore the size, press the "ESC" key

Refresh Browser Window

If the webcast freezes and does not recover in 3-4 seconds, refresh browser window

Freezing or Stuttering Issues

Adjust the amount of bandwidth needed by putting your mouse anywhere over the video window. A tool bar will appear at the bottom. On the right side you will see a "HD" button, click on that button and you will see a list of options. The top is "auto", with decreasing numbers below. Select a lower bandwidth (such as 360p) to see if your webcast improves

For Technical Support call

1-800-504-5379

Mobile Viewing Requirements

Android Devices

Android 2.3+ with Adobe Flash Player 10.2 or better installed
[Install Flash Player](#)

Apple Devices

iOS 4+

Online Viewing Requirements

Bandwidth

512kbps

Required Hardware and Software	Screen resolution of 1024X768 or larger Sound card and speakers/headphones
Browser	Microsoft Internet Explorer 7.0 or better Mozilla Firefox 4 or better Safari 5 or better
Windows	Operating System: Windows 8 desktop mode, Windows 7; Windows Vista; Windows XP Service Pack 2 or 3 x86 or x64 (Browsers must be in 32-bit mode) 1.6-gigahertz (GHz) or higher processor 512MB of RAM
Mac OS	Operating System: Apple Mac OS X 10.4.8 or above Intel Core™ Duo 1.83GHz or faster processor 512MB of RAM

Registration and Fees

ondemand.acmg.net

ACMG Members and ACMG Trainees: (\$50)

Non-members (\$60)

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Questions regarding CE credit should be directed to education@acmg.net.

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