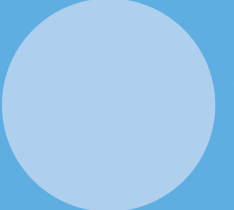


Cardiology in the Young

December 2014 HeartWeek Issue of Cardiology in the Young



Johns Hopkins All Children's Heart Institute

14th International Symposium on Congenital Heart Disease Special Focus: Diseases of the Cardiac Valves from the Fetus to the Adult Co-Sponsor: The American Association for Thoracic Surgery (AATS) Renaissance Vinoy Resort & Golf Club, Saint Petersburg, Florida February 15-18. 2014

and

HeartWeek 2014:

The Children's Hospital of Philadelphia Cardiology 2014: 17th Annual Update on Pediatric

INNOVATIONS: What We Know.

What We Need to Learn. Where We Need to Be. Disney's Yacht and Beach Club Resorts, Lake Buena Vista, Fla. February 19 – 23, 2014





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Cardiology in the Young is indexed and abstracted in Index Medicus/MEDLINE; Current Contents/Clinical Medicine; Research Alert; Sci Search; EMBASE/Excerpta Medica

Commissioning Editor: Dan Edwards Production Editor: Nicki Marshall

Design and Production: MPS Limited, Bangalore Printed & Bound: The Sheridan Press, USA

Published by Cambridge University Press (Journals), UPH, Shaftesbury Road, Cambridge CB2 8BS, UK Tel: +44 (0)1223 326491; Fax: +44 (0)1223 325802; E-mail: dedwards@cambridge.org

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DECEMBER 2014 HEARTWEEK ISSUE OF CARDIOLOGY IN THE YOUNG

HeartWeek 2014

Highlights of HeartWeek 2014: Diseases of the Cardiac Valves from the Foetus to the Adult

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This issue was supported by: Johns Hopkins All Children's Heart Institute, Saint Petersburg, Florida



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Jacobs JP, Quintessenza JA, Dadlani G, Everett AD, Anderson RH (Editors). December 2014 Issue of Cardiology in the Young: Annual Heart Week in Florida Issue Number 12 – **Highlights of HeartWeek 2014: Diseases of the Cardiac Valves from the Foetus to the Adult**. Cardiology in the Young, Volume 24, Issue 6, pages 959–1153, December, 2014.

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The International Society for Nomenclature of Paediatric and Congenital Heart Disease

■HE INTERNATIONAL SOCIETY FOR NOMENCLATURE of Paediatric and Congenital Heart Disease (ISNPCHD) is an established not-for-profit organization, incorporated in Canada, with the collective mission to identify, standardize, and maintain an international system of nomenclature, the International Paediatric and Congenital Cardiac Code (IPCCC), to enhance global communication and facilitate patient care, research, and training in paediatric and congenital heart care across disciplines. In other words, the IPCCC provides a common language and terminology, inclusive of definitions and imaging, which covers the entire field of diagnostic and procedural terms for paediatric and congenital heart care. The IPCCC is the product of the cross-mapping work by the ISNPCHD Nomenclature Working Group during the decade following the publication in 2000 of two similar and complementary nomenclature systems, namely the International Congenital Heart Surgery Nomenclature and Database Project under the auspices of the European Association for Cardio-Thoracic Surgery and Society of Thoracic Surgeons, and the European Paediatric Cardiac Code under the auspices of the Association for European Paediatric Cardiology. The IPCCC is owned by the ISNPCHD but is digitally published for free download for private use (www.ipccc. net) in these two Societal versions, both with a Long List of over 10,000 terms with qualifiers, and a Short List of up to 1000 terms for use in databases when comparing institutional outcomes, both nationally and internationally. The Long Lists of the IPCCC comprehensively cover the field of paediatric and congenital cardiac care, including diagnoses of congenital and related acquired pathology, comorbid conditions, transcatheter and operative procedures, and a full list of postprocedural complications. The Short Lists are used within databases across the world with over 500,000 registered patients. The ISNPCHD encourages the commercial use of the

IPCCC by requiring and providing free-of-charge license agreements for its use, to ensure that the IPCCC remains unaltered by parties other than the ISNPCHD.

Since 2007 the ISNPCHD has been operating through three working groups:

- The Nomenclature Working Group, which continues to maintain, develop, expand, update, and preserve the IPCCC;
- The Definitions Working Group, which is engaged in writing definitions for the terms in the IPCCC. More recently this initiative has focused on the terms provided by the ISNPCHD at the behest of the World Health Organization (WHO) for the 11th revision of the International Classification of Diseases;
- The Archiving Working Group, which is engaged in linking images and videos to the IPCCC, including cardiac morphologic specimens, echocardiography, angiography, computerized axial tomography, magnetic resonance imaging, intraoperative photographs and intraoperative videos.

The ISNPCHD, through the IPCCC, enables institutions from around the world to seamlessly communicate with each other, comparing and then improving outcomes and the quality of care that is given to children, young people, and adults born with malformed hearts. This common language enables institutions to learn from those hospitals performing best at a global level, as well as facilitating research projects, such as comparing the longer term quality of life and complications in those who have required operative and transcatheter interventions. In addition, the nomenclature with corresponding definitions and matching imaging, enhances teaching of this specialty to the next generation of clinicians dedicated to pediatric and congenital cardiac care, both in the developed and developing world.





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ISTANBUL, TURKEY











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Cardiology in the Young © 2014 Cambridge University Press, ISSN 1047-9511 is published bi-monthly

Subscription information: Volume 24 (6 issues) will appear in 2014. Correspondence concerning subscriptions should be addressed to: Journals Customer Services, Cambridge University Press, UPH, Cambridge CB2 8BS, UK. Tel: +44 (0)1223 325150; E-mail enquiries: journals_subscriptions@cambridge.org; Web: www.cambridge.org; Cambridge University Press (Journals Online: www.journals.cambridge.org). The submission of a manuscript implies the following: (a) that the work described has not been published before, except in the form of an abstract or as part of a published lecture, review, or thesis; (b) that it is not under consideration for publication elsewhere; (c) that all co-authors approve its publication; (d) that its publication is approved by the responsible authorities at the institute where the work has been carried out; (e) that when the manuscript is accepted for publication, the authors agree to the automatic transfer of the copyright to the publisher; (f) that the manuscript will not be published elsewhere in any language without the consent of the copyright holders, and (g) that written permission of the toopyright by the authors for material used from other copyrighted sources. All articles published in this journal are protected by copyright, which covers the exclusive rights to reproduce and distribute the article, as well as all translation rights. No material published in this journal may be reproduced photographically or stored on microfilm, in video disks, etc., without first obtaining written permission from the publishers. The use of general descriptive names, trade names, trademarks, etc., in this publication, even if not specifically identified, does not imply that these names are not protected by the relevant laws and regulations. While the advice and information in this journal are believed to be true and accurate at the date of its going to press, neither the authors, the editors, the publisher, nor any affiliated Society can accept any legal responsibility for any errors or ormissions that m