

This is a repository copy of National clinical audit data decodes the genetic architecture of developmental dysplasia of the hip.

White Rose Research Online URL for this paper: <a href="https://eprints.whiterose.ac.uk/120862/">https://eprints.whiterose.ac.uk/120862/</a>

Version: Accepted Version

# **Proceedings Paper:**

Hatzikotoulas, K., Roposch, A., Shah, K.M. et al. (13 more authors) (2017) National clinical audit data decodes the genetic architecture of developmental dysplasia of the hip. In: Osteoarthritis and Cartilage. 2017 OARSI World Congress on Osteoarthritis: Promoting Clinical and Basic Research in Osteoarthritis, 27-30 Apr 2017, Las Vegas, USA. Elsevier, S36-S37.

https://doi.org/10.1016/j.joca.2017.02.073

Article available under the terms of the CC-BY-NC-ND licence (https://creativecommons.org/licenses/by-nc-nd/4.0/).

### Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs (CC BY-NC-ND) licence. This licence only allows you to download this work and share it with others as long as you credit the authors, but you can't change the article in any way or use it commercially. More information and the full terms of the licence here: https://creativecommons.org/licenses/

## Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



Print this Page for Your Records

Close Window

Control/Tracking Number: 17-A-255-OARSI **Activity:** Abstract Current Date/Time: 11/21/2016 2:26:29 AM

#### NATIONAL CLINICAL AUDIT DATA DECODES THE GENETIC ARCHITECTURE OF DEVELOPMENTAL DYSPLASIA OF THE HIP

Author Block: K. Hatzikotoulas<sup>1</sup>, A. Roposch<sup>2</sup>, K. M. Shah<sup>3</sup>, M. J. Clark<sup>3</sup>, S. Bratherton<sup>3</sup>, V. Limbani<sup>4</sup>, K. Warsame<sup>2</sup>, M. Ratnayake<sup>5</sup>, M. Tselepi<sup>5</sup>, J. Schwartzentruber<sup>1</sup>, J. Steinberg<sup>1</sup>, D. DH Case Control Consortium<sup>2</sup>, J. Loughlin<sup>5</sup>, D. M. Eastwood<sup>4</sup>, E. Zeggini<sup>1</sup>, **J. M. Wilkinson**<sup>3</sup>; <sup>1</sup>Wellcome Trust Sanger Inst., Cambridge, United Kingdom, <sup>2</sup>Univ. Coll. London, London, United Kingdom, <sup>3</sup>Univ. of Sheffield, Sheffield, United Kingdom, <sup>4</sup>Royal Natl. Orthopaedic Hosp., Stanmore, United Kingdom, <sup>5</sup>Newcastle Univ., Newcastle, United Kingdom

#### **Abstract:**

Purpose: Developmental dysplasia of the hip (DDH) is a heritable condition with an incidence of 3.6 per 1000 live births in the United Kingdom. DDH is characterised by abnormal development of the hip joint that results in pain, loss of function, and secondary osteoarthritis. We applied case identification using national clinical audit data and postal recruitment to conduct the first successful genome-wide study of the genetic architecture of DDH to better understand its biological aetiology.

Methods: We recruited 770 patients (639 female) with a history of DDH from the English National Joint Registry (NJR) and 3364 controls (3048 female) from UK Household Longitudinal Study (UKHLS) for the discovery cohort. All participants were of UK European ancestry. Genomic DNA was genotyped using the Illumina HumanCoreExome beadchip. Following quality control checks at the sample and genotype level, association analyses were conducted under an additive model. Identified independent signals were followed up in an independent replication cohort of 1129 (1004 female) children with DDH, recruited prospectively, and 4652 independent controls from the UKHLS. Finally, a meta-analysis of both cohorts was conducted. We defined genome-wide significance as p<5x10<sup>-8</sup>. We

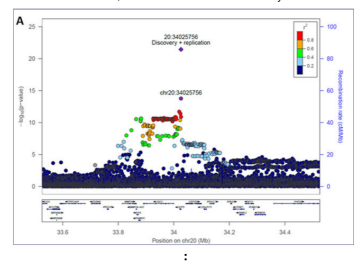
estimated the heritability of DDH using genetic complex trait analysis (GCTA). To test for shared genetics of DDH with OA, we employed high-resolution polygenic risk scoring (with data from a previous GWAS of OA) by using evenly spaced p-value thresholds between 0.001 and 0.50.

Results: Using genome-wide single nucleotide polymorphism (SNP) data and GCTA analysis we find that commonfrequency autosomal SNPs explain 55% (±se=6%, p<0.0001) of the liability-scale heritability of DDH. In the discovery case-control analysis we find 53 SNPs, comprising 25 independent signals, showed suggestive association with DDH at p<9x10<sup>-5</sup>. Eleven correlated variants reached genome wide significance, with rs 143384 in the GDF5 promoter as the lead variant (OR 1.57, 95% CI 1.3-1.77, p=1.72x10<sup>-14</sup>). At replication, the rs143384 variant was also associated with DDH at genome-wide significance (OR=1.37, 95% CI 1.24 to 1.51, p=1.33x10<sup>-10</sup>). Finally, at meta-analysis the rs143384 variant was associated with DDH with OR=1.44 (95% CI 1.34 to 1.56, p=3.55x10<sup>-22</sup>, Figure A; regional association plot showing variants within GDF5 locus and strength of association). We also identify two further replicating loci with suggestive association to DDH near the NFIB (rs4740554, OR 1.30 [95% CI 1.16-1.45], p= $4.44\times10^{-6}$ ) and LOXL4 (rs4919218, 1.19 [1.10-1.28] p= $4.38\times10^{-6}$ ) genes. We identify RETSAT and PDRG1 association with DDH through gene-based analysis (p=1·19x10<sup>-6</sup> and p=3·77x10<sup>-6</sup>, respectively). To identify potential shared genetic aetiology and hence common biological pathways underpinning DDH and idiopathic hip OA we

constructed polygenic risk scores using the arcOGEN dataset and tested their predictive potential in the DDH GWAS. We find no significant association between polygenic risk scores for hip OA and DDH.

Conclusions: Using the NJR as a proof-of-principle, we describe the genetic architecture of DDH and establish the first robust DDH genetic locus. Fine mapping of the 5'UTR of GDF5 indicates rs143384 (rather than 143383) as the causal variant. We also demonstrate a robust and scalable national clinical audit-based recruitment strategy for genetic studies that is transferrable to other complex diseases.

> Table 1: Replicating variants associated with DDH



Category (Complete): Genetics and Genomics and Epigenetics Keyword (Complete): Genetics; Hip; Outcome Presentation Preference (Complete): Either Podium or Poster Additional (Complete):

Type of Abstract (Required): Clinical Science Are you age 40 or under and within 5 years of your research degree? (Required): No Do you want to be considered for a Need Based Travel Assistance Award? (Required): No

Payment (Complete): Your credit card order has been processed on Monday 21 November 2016 at 2:25 AM.

Status: Complete

Osteoarthritis Research Society International 1120 Rt. 73, Ste. 200 Mt. Laurel, NJ 08054, USA vconverse@oarsi.org

For any technical inquiries, <u>click here</u> to contact OASIS Helpdesk or call 217-398-1792.

**Leave OASIS Feedback** 

Powered by <u>cOASIS</u>, The Online Abstract Submission and Invitation System <sup>SM</sup> © 1996 - 2016 CTI Meeting Technology All rights reserved.