

Publication / Bibliography List

Roger A. Hubbard, PhD - Journal Articles

1. Frantz C, DM Sekora, DC Henley, CK Huang, Q Pan, NB Quigley, E Gorman, RA Hubbard and I Mirza (2007). Comparative valuation of three JAK2V617F mutation detection methods. *American Journal of Clinical Pathology*. 128: 865-874.
2. Hubbard RA (2003). Human papillomavirus testing methods. *Archives Pathology Laboratory Medicine* 127:940.
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4. Modarress KJ, Cullen AP, Jaffurs WJ, Troutman GL, Mousavi N, Hubbard RA, Henderson S, Lorincz AL (1999). Detection of Chlamydia trachomatis and Neisseria gonorrhoeae in swab specimens by the Hybrid Capture II and PACE 2 nucleic acid probe tests. *Sexually Transmitted Diseases* 26:303.

Ron Lee, MD - Journal Articles

1. Lee, R.V., Green C.A., Negrea O.G., Dodson S., Farrell S.K., Hewitt J.E., Jago T., Ramsey C.E., Cato T.C., Crawford E.C., Henley D.C., Phelan M.C., Potter N.T. (2009). B-cell lymphoma with intermediate- to high-grade features and different immunophenotypic profiles involving separate anatomic sites with a good response to R-CHOP. *LabMedicine*. 40(2):79-86.
2. Davis BH et al. (2007) Bethesda International Consensus Recommendations on the Flow Cytometric Immunophenotypic Analysis of Hematolymphoid Neoplasia: Medical Indications, pp. S5-S13 (Cytometry, Volume 72B, Supplement 1, 27 Aug. 2007)
3. Lee RV (2006). Comparison of Inbound and Outbound Diagnoses for Samples Submitted for Flow Cytometric Analysis. (*LabMedicine*, 2007 Apr. 38(4):240-243)

Ron Lee, MD - Abstracts/Posters/Presentations

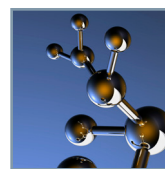
1. Lee RV, Braylan R (June 2002). CBC, manual differential, and flow cytometric analyses of whole blood samples from normal volunteers stored at room and cold temperatures. Abstract/Resident Research Day presentation, University of Florida.

Katy Phelan, PhD, FACMG - Journal Articles

1. Lee, R.V., Green C.A., Negrea O.G., Dodson S., Farrell S.K., Hewitt J.E., Jago T., Ramsey C.E., Cato T.C., Crawford E.C., Henley D.C., Phelan M.C., Potter N.T. (2009). B-cell lymphoma with intermediate- to high-grade features and different immunophenotypic profiles involving separate anatomic sites with a good response to R-CHOP. *LabMedicine*. Feb;40(2):79-86.
2. Wilson HL, Wong ACC, Tse WY, Stapleton GA, Phelan MC, McDerimid HE (2003). Molecular characterization of the 22q13 deletion syndrome supports the role of haploinsufficiency of SHANK3/PROSAP2 in the major neurological symptoms. *Journal of Medical Genetics* 40:575.
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7. Phelan MC, Saul RA, Gailey TA Jr, Skinner SA (1995). Prenatal diagnosis of mosaic 4p- in a fetus with trisomy 21. *Prenatal Diagnosis* 15:274.
8. Shapiro LR, Simensen RJ, Wilmot PL, Fisch GS, Vibert BK, Fenwick RG, Tarleton J, Phelan MC (1994). Asymmetry of methylation with FMR-a full mutation in two 45,X/45,XX mosaic females associated with normal intellect. *American Journal Medical Genetic* 51:507.
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10. McConkie-Rosell A, Lachiewicz AM, Spiridigliozzi GA, Tarleton J, Schoenwald S, Phelan MC, Goonewardena P, Ding X, Brown WT (1993). Evidence that methylation of the FMR-1 locus is responsible for variable phenotypic expression of the Fragile X syndrome. *American Journal Medical Genetic* 53:800.
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Katy Phelan, PhD, FACMG - Books/Chapters

1. Phelan K. 22q13.3 deletion syndrome. In: *GeneReviews*, <http://www.genetests.org>, in press.
2. Phelan MC, Hall JG. Twins. In: *Human Malformations and Related Anomalies*, Vol II, 2nd edition, RE Stevenson and JG Hall, eds., Oxford University Press, New York, in press.
3. Phelan MC, Stapleton GA, Rogers RC (2005). 22q13 deletion syndrome. In: *Management of Genetic Syndromes*, 2nd edition, Cassidy SB, Allanson JE, eds., John Wiley & Sons, Hoboken, NJ.
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Nicholas T. Potter, PhD, FACMG - Journal Articles

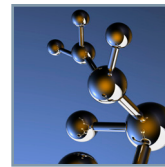
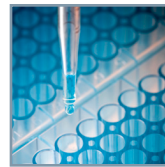
1. Lee, R.V., Green C.A., Negrea O.G., Dodson S., Farrell S.K., Hewitt J.E., Jago T., Ramsey C.E., Cato T.C., Crawford E.C., Henley D.C., Phelan M.C., Potter N.T. (2009). B-cell lymphoma with intermediate- to high-grade features and different immunophenotypic profiles involving separate anatomic sites with a good response to R-CHOP. *LabMedicine*. Feb;40(2):79-86.
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Nicholas T. Potter, PhD, FACMG - Books/Chapters

1. Potter, NT (2008) Standards and recommendations for molecular diagnostic testing for Huntington disease, the autosomal dominant spinocerebellar ataxias, and Friedreich ataxia. In: *Handbook of Molecular Diagnostics*, WW Grody et al., eds., Elsevier, San Diego CA in press.
2. Potter NT (2006). Neurodegenerative Disorders. In: *Molecular Pathology in Clinical Practice*, DGB Leonard, A Bagg, A Caliendo, K Kaul, K Snow-Bailey, and V Van Deerlin eds., Springer-Verlag, New York, NY, pp.179-190.
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Nicholas T. Potter, PhD, FACMG - Abstracts/Posters/Presentations

1. Jago T, Morgan M, Hubbard RA, Potter NT (2006). Evaluation of the Gen-Probe APTIMA Combo 2 (AC2) Assay and the TIGRIS DTS Automated Workstation for the Detection of CT and NG rRNA from Liquid Cytology (ThinPrep® and SurePath™) Specimens. *Journal of Molecular Diagnostics* 8:643.
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Neil B. Quigley, PhD, HCLD(ABB) - Journal Articles

1. Jones D, Kamel-Reid S, Bahler D, Doug H, Elenitoba-Johnson K, Press R, Quigley N, Rothberg P, Sabath D, Viswannatha D, Weck K, Kehnder J. (2009). Laboratory practice guidelines for detecting and reporting BCR-ABL drug resistance mutations in chronic myelogenous leukemia and acute lymphoblastic leukemia. *Journal of Molecular Diagnostics*. Jan; 11:4-11.
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Neil B. Quigley, PhD, HCLD(ABB) - Abstracts/Posters/Presentations

1. Day, SP, NB Quigley, DI Quigley, B Yen-Lieberman and M Olson (2008). Analytical performance characteristics of the CE-marked Invader[®] HPV HR molecular assay. HPV in Human Pathology Congress (Prague, Czech Republic; May 1-3).
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3. Henley, DC and NB Quigley (2007). Validation of a novel quantitative PCR test for the JAK2V617F mutation commonly associated with clonal myeloproliferative disease. (SE Regional Group Meeting of the Association of Genetics Technologists, Knoxville, Tennessee; September).
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L. Rane Taylor, PhD - Journal Articles

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Elizabeth Stone, BS, MT (ASCP) - Clinical and Laboratory Guidelines

1. Stetler-Stevenson M, Ahmad E, Barnett D, Braylan RC, DiGiuseppe JA, Marti G, Menozzi D, Oldaker TA, Orfao de Matos A, Rabellino E, Stone EC, Walker C (2007). *Clinical Flow Cytometric Analysis of Neoplastic Hematolymphoid Cells; Approved Guidelines-Second Edition*. CLSI document H43-A2.