

**CURRICULUM VITAE****INFORMAZIONI PERSONALI**

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|-----------------|-------------------------------------|
| Cognome Nome    | <b>Acquila Maura</b>                |
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| Nazionalità     | italiana                            |
| Data di nascita | 09/09/1959                          |

**FORMAZIONE E STAGE**

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| Tipologia |  |
| Data      |  |
| Sede      |  |

**SPECIALIZZAZIONI**

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| Tipologia | Genetica medica |
| Data      | Ottobre 1994    |
| Sede      | Genova          |

**ESPERIENZA LAVORATIVA**

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| Data (da – a)      | 1987 a tutt'oggi   |
| Nome Istituzione   | istituto G. Gaslini laboratorio ematologia ed emofilia IV pediatria                          |
| Incarico ricoperto | Dal 1987 al 1993 Biologa borsista; dal 15 marzo 1993 dirigente biologa a tempo indeterminato |

**COORDINAMENTO GRUPPI DI LAVORO – GRUPPI DI RICERCA. NAZIONALI ED INTERNAZIONALI**

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| <b>ELENCO DELLE PUBBLICAZIONI SCIENTIFICHE PIÙ SIGNIFICATIVE</b> | <p>Morerio C, ACQUILA M, Rosanda C, Rapella A, Dufour C, Locatelli F, Maserati E, Pasquali F, Panarello C. HCMOGT-I is a novel fusion partner to PDGFRB in juvenile myelomonocytic leukemia with t(5;17)(q33;pl1.2). <i>Cancer Res</i> 2004;64:2649-51.</p> <p>ACQUILA M, Pasino M, Lanza T, Bottini F, Molinari AC, Bicocchi MP. Duplication of exon 13 causes one third of the cases of mild hemophilia A in northern Italy. <i>Haematologica</i> 2004;89:758-9.</p> <p>ACQUILA M, Pasino M, Lanza T; Molinari ACM, Caprino D, Bottini F, Bicocchi MP. Identification of mutations in exon 14 including 5 novelties in 13 Italian patients with haemophilia A. <i>Haemophilia</i> 2004;10:744-6.</p> <p>Bicocchi MP, Pasino M, Lanza T, Bottini F, Molinari AC, Caprino D, Rosano C, ACQUILA M. Small FVIII gene rearrangements in 18 Hemophilia A patients: five novel mutations. <i>Am J Haematol</i> 2005;78:117-22.</p> <p>Bicocchi MP, Migeon BR, Pasino M, Lanza T, Bottini F, Boeri E, Molinari AC, Corsolini F, Morerio C, ACQUILA M. Familial nonrandom inactivation linked to the X inactivation centre in heterozygotes manifesting haemophilia A. <i>Eur J Hum Genet</i> 2005;13: 635-40.</p> <p>Bicocchi MP., Pasino M, Lanza T, Bottini F, Molinari AC, Rosano C, ACQUILA M. Ectopic mRNA analysis and molecular modelling substantiate severe haemophilia in a patient with a FVIII gene splice mutation. <i>Thromb Haemost</i> 2005;93:391-2.</p> <p>Morerio C, ACQUILA M., Rosanda C, Rapella A, Tassano E, Micalizzi C, Panarello C. t(9;11)(p22;p15) with NUP98-LEDGF fusion gene in pediatric acute myeloid leukemia. <i>Leuk Res</i> 2005;29:467-70.</p> <p>Bicocchi MP, Pasino M, Rosano C, Molinari AC, Della VE, Lanza T, et al. Insight into molecular changes of the FIX protein in a series of Italian patients with haemophilia B. <i>Haemophilia</i> 2006 May;12(3):263-70.</p> <p>Morerio C, Acquila M, Rapella A, Tassano E, Rosanda C, Panarello C. Inversion (11)(p15q22) with NUP98-DDX10 fusion gene in pediatric acute myeloid leukemia. <i>Cancer Genet Cytogenet</i> 2006 Dec;171(2):122-5.</p> <p>Acquila M, Pasino M, Di DM, Bottini F, Molinari AC, Bicocchi MP. MLPA assay in F8 gene mutation screening. <i>Haemophilia</i> 2008 May;14(3):625-7.</p> <p>Fabiano C, Acquila M, Bicocchi MP, Sammarco P. An uncommon case of a female carrier of two distinct X-linked disorders. <i>Haemophilia</i> 2008 May;14(3):665-6.</p> <p>Santacroce R, Acquila M, Belvini D, Castaldo G, Garagiola I, Giacomelli SH, et al. Identification of 217 unreported mutations in the F8 gene in a group of 1,410 unselected Italian patients with hemophilia A. <i>J Hum Genet</i> 2008;53(3):275-84.</p> |
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| ULTERIORI INFORMAZIONI |  |
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