

CURRICULUM VITAE

Dott. MARIA CRISTINA DIGILIO

settembre 2008

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Data di nascita: 14 maggio 1961

Stato civile: nubile

Indirizzo permanente: Viale Marco Polo 77, 00154 Roma. Tel. 06/5743229

Indirizzo ospedale: Ospedale Pediatrico Bambino Gesù, U.O. Genetica Medica, Piazza S.Onofrio 4, 00165, Roma. Tel. 06/68592227

TITOLI DI STUDIO:

- Laurea in Medicina e Chirurgia, presso l'Università degli Studi di Roma La Sapienza, 24/10/86
- Abilitazione all'esercizio della professione medico-chirurgica, presso l'Università degli Studi La Sapienza di Roma, Novembre 86
- Specializzazione in Pediatria, presso l'Università degli Studi La Sapienza di Roma, 27/6/90
- Specializzazione in Genetica Medica, presso l'Università Cattolica del Sacro Cuore di Roma, 9/6/95
- Iscrizione all'Ordine dei Medici Chirurghi di Roma, dal 2/4/87 a tutt'oggi

ATTIVITA' PROFESSIONALE

- Servizio in qualità di Dirigente Medico di I livello presso l'U.O. di Genetica Medica dell'Ospedale Bambino Gesù di Roma: dal 5/8/94 a tutt'oggi
- Responsabile dell'Alta Specializzazione in Dismorfologia presso lo stesso ospedale: dall' 1/1/2002 a tutt'oggi

STAGES DI FORMAZIONE ALL'ESTERO

- Stage presso il Baltimore-Washington Infant Study Group, University of Maryland, Baltimore, USA, novembre 2002

ATTIVITA' DIDATTICA

- Corso Universitario per Infermieri, Università di Tor Vergata, Roma, Anni accademici: 2000-2001 a tutt'oggi
- Corsi di Scuola Medica Ospedaliera "Sindromi genetiche in Pediatria: Clinica e Laboratorio", Anni accademici: 2002-2003, 2005-2006.

ISCRIZIONE A SOCIETA' MEDICHE

- Ordine dei Medici di Roma e Provincia
- Società Italiana di Genetica Umana (SIGU)
- Società Italiana di Pediatria (SIP)
- SIMGePeD
- Società Italiana di Endocrinologia Pediatrica (SIEDP)

REVISORE DELLE SEGUENTI RIVISTE

- American Journal of Medical Genetics
- Clinical Genetics
- American Journal of Cardiology
- European Journal of Echocardiography

LETTURE / CONFERENZE TENUTE ALL'ESTERO SU INVITO

- International Meeting on the etiology and morphogenesis of congenital heart diseases, Tokyo, Japan, November 7-9, 2002.
- 12th International Scientific Congress on the DiGeorge syndrome, Strasbourg, France, July 7-9, 2006
- The fifth International 22q11.2 deletion syndrome Conference, Marseille, France, July 10-11, 2006
- Rare disorders of the MAPK pathway: Current status / future directions. Barcelona, Spain, May 30-31, 2008

RELATORE SU INVITO in circa 25 meetings e congressi italiani

ARTICOLI IN GIORNALI INDEX

2008

Digilio MC, Calzolari F, Capolino R, Toscano A, Sarkozy A, de Zorzi A, Dallapiccola B, Marino B. Congenital heart defects in patients with oculo-auriculo-vertebral spectrum (Goldenhar syndrome). *Am J Med Genet* 2008;146A:1815-1819.

Carotti A, **Digilio MC**, Piacentini G, Saffirio C, Di Donato RM, Marino B. Cardiac defects and results of cardiac surgery in 22q11.2 deletion syndrome. *Dev Disabil Res Rev* 2008;14:35-42.

Digilio MC, Sarkozy A, Capolino R, Chiarini Testa MB, Esposito G, de Zorzi A, Cutrera R, Marino B, Dallapiccola B. Costello syndrome: clinical diagnosis in the first year of life. *Eur J Pediatr* 2008;167:621-628.

Sarkozy A, **Digilio MC**, Dallapiccola B. LEOPARD syndrome. *Orphanet J Rare Dis* 2008; 3: 13.

Digilio MC, Marino B, Dallapiccola B. Deletion 22q11 and isolated congenital heart disease. *Int J Cardiol* 2008; 123:364-365.

Limongelli G, Sarkozy A, Pacileo G, Calabrò P, **Digilio MC**, Maddaloni V, Gagliardi G, Di Salvo G, Iacomino M, Marino B, Dallapiccola B, Calabrò R. Genotype-phenotype analysis and natural history of left ventricular hypertrophy in LEOPARD syndrome. *Am J Med Genet* 2008;146A:620-628.

Digilio MC, Capolino R, Dallapiccola B. Autosomal dominant transmission of nonsyndromic diastasis recti and weakness of the linea alba. *Am J Med Genet* 2008;146A:254-256.

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Limongelli G, Pacileo G, Melis D, Calabrò P, **Digilio MC**, Sarkozy A, Maddaloni V, Capozzi G, Sebastio G, Andria G, Calabrò R. Trisomy 18 and hypertrophic cardiomyopathy in an 18-year-old woman. *Am J Med Genet* 2008;146A:327-329.

2007

Pandit B, Sarkozy A, Pennacchio LA, Carta C, Oishi K, Martinelli S, Pogna EA, Schackwitz W, Ustaszewska A, Landstrom A, Bos JM, Ommen SR, Esposito G, Lepri F, Faul C, Mundel P, Lopez Sigüero JP, Tenconi R, Selicorni A, Rossi C, Mazzanti L, Torrente I, Marino B, **Digilio MC**, Zampino G, Ackerman MJ, Dallapiccola B, Tartaglia M, Gelb BD. Gain-of-function RAF1 mutations cause Noonan syndrome and LEOPARD syndromes with hypertrophic cardiomyopathy. *Nat Genet* 2007;8:1007-1012.

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Limongelli G, Pacileo G, **Digilio MC**, Calabrò P, Di Salvo G, Rea A, Miele T, Frigiola A, Sarkozy A, Dallapiccola B, Marino B, Calabrò R. Severe, obstructive biventricular hypertrophy in a patient with Costello syndrome: Clinical impact and management. *Int J Cardiol* 2007;Epub ahead of print.

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2006

Digilio MC, Sarkozy A, de Zorzi A, Pacileo G, Limongelli G, Mingarelli R, Calabrò R, Marino B, Dallapiccola B. LEOPARD syndrome: clinical diagnosis in the first year of life. *Am J Med Genet A* 2006;140: 740-746.

Digilio MC, Sarkozy A, Pacileo G, Limongelli G, Marino B, Dallapiccola B. PTPN11 gene mutations: linking the Gln510Glu mutation to the "LEOPARD syndrome phenotype". *Eur J Pediatr* 2006;165:803-5.

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2005

Digilio MC, Marino B, Capolino R, Angioni A, Sarkozy A, Roberti MC, Conti E, de Zorzi A, Dallapiccola B. Familial recurrence of nonsyndromic congenital heart defects in

first degree relatives of patients with deletion 22q11.2. *Am J Med Genet* 2005;134:158-164.

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2004

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2003

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2002

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2001

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2000

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1999

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