

N° 60 PUBBLICAZIONI SU RIVISTE INTERNAZIONALI :

- 1: Cinzia Ciccacci, Mattia Falconi, Nicoletta Paolillo, Francesco Oteri, Vittorio Forte, Giuseppe Novelli, Alessandro Desideri and Paola Borgiani
Characterization of a novel CYP2C9 gene mutation and structural bioinformatic protein analysis in a warfarin hyper-sensitive patient
Pharmacogenetics and Genomics, IN PRESS 2011
- 2: C Ciccacci, N Paolillo, D Di Fusco, G Novelli and P Borgiani
EPHX1 polymorphisms are not associated with warfarin response in an Italian population.
Clinical Pharmacology & Therapeutics, IN PRESS 2011
- 3: Saccucci P, Meloni GF, Verrotti A, Borgiani P, D'Annibale F, Giannini C, Lucarelli P, Bottini N, Chiarelli F, Bottini E, Gloria-Bottini F.
A study of three polymorphic sites of the ADA gene in children with type 1 diabetes mellitus.
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ATG16L1 Ala197Thr is not associated with susceptibility to Crohn's disease or with phenotype in an Italian population.
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- 11: Borgiani P, Perricone C, Ciccacci C, Romano S, Novelli G, Biancone L, Petruzzello C, Pallone F.
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