



MASTER PEDIATRIC ENDOCRINOLOGY AND DIABETES

Sabrina Giglio list of publications

Stagi S, Ricci F, Bianconi M, Sammarco MA, Municchi G, **Toni S**, Lenzi L, Verrotti A, de Martino M. Retrospective Evaluation of Metformin and/or Metformin Plus a New Polysaccharide Complex in Treating Severe Hyperinsulinism and Insulin Resistance in Obese Children and Adolescents with Metabolic Syndrome. *Nutrients*. 2017 May 20;9(5). pii: E524. doi: 10.3390/nu9050524. [[ABSTRACT](#)] [[FULL TEXT](#)] [[PDF](#)]

Galluzzi F, Stagi S, Salti R, **Toni S**, Piscitelli E, Simonini G, Falcini F, Chiarelli F. [Osteoprotegerin serum levels in children with type 1 diabetes: a potential modulating role in bone status](#). *Eur J Endocrinol*. 2005 Dec;153(6):879-85. [[ABSTRACT](#)] [[FULL TEXT](#)] [[PDF](#)]

Bassi A, **Toni S**, Piccini B, Giacalone M, DE Martino M. Yellow skin discoloration in a 4 years-old girl with type 1 diabetes mellitus. *G Ital Dermatol Venereol*. 2017 Nov 30. doi: 10.23736/S0392-0488.17.05790-X. [[ABSTRACT](#)]

Arcangeli A, Lastraioli E, Piccini B, D'Amico M, Lenzi L, Pillozzi S, Calabrese M, **Toni S**, Arcangeli A. Circulating Endothelial Progenitor Cells in Type 1 Diabetic Patients: Relation with Patients' Age and Disease Duration. *Front Endocrinol (Lausanne)*. 2017 Oct 23;8:278. doi: 10.3389/fendo.2017.00278. [[ABSTRACT](#)] [[FULL TEXT](#)] [[PDF](#)]

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Scaramuzza AE, Arnaldi C, Cherubini V, Piccinno E, Rabbone I, **Toni S**, Tumini S, Candela G, Cipriano P, Ferrito L, Lenzi L, Tinti D, Cohen O, Lombardo F. Recommendations for the use of sensor-augmented pumps with predictive low-glucose

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Curriculum Vitae

Author list publications

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