

# Stefano Volpi, MD, PhD

Reumatologia e Malattie Autoinfiammatorie  
IRCCS Istituto Giannina Gaslini, Genova, Italy

Nato il 17/10/1980 a Genova

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## Track Record

Scientific output, ottenuto da Scopus il 25/07/2024

Numero totali di articoli pubblicati su riviste indicizzate: 136

Citazioni: 5853

h-index: 40

Articoli come primo autore: 7

Articoli come ultimo autore o autore corrispondente: 4

## Biosketch

Nella mia carriera mi sono concentrato sullo studio delle cause genetiche delle immunodeficienze e sul meccanismo di queste malattie attraverso modelli in vitro tramite cellule staminali pluripotenti indotte e modelli animali.

Durante la mia specializzazione e dottorato di ricerca (2007-2013) nell'ambito della ricerca svolta ad Harvard, negli Stati Uniti, ho descritto come primo autore una nuova immunodeficienza causata dalla mutazione nel gene EXTL3 (JEM PMID 28148688). Ho contribuito alla descrizione di altre 3 immunodeficienze (Cell PMID 29474921, Nature PMID 25307056 e Science PMID 25814066). Ho eseguito studi scientifici di base sulle immunodeficienze primarie utilizzando modelli murini o iPSC (Blood PMID 22302739; Nature Immunology **Journal cover** PMID 30127432; JACI come primo autore PMID 26409660; PNAS PMID 30154162; Frontiers in Immunology PMID 25101082).

Mi sono trasferito a Losanna nel 2014 dove ho allestito da zero il nuovo laboratorio del mio nuovo PI e ho completato uno studio su un modello animale come primo autore (Blood **Journal cover** PMID 26468226).

Tornato in Italia nel 2015, mi sono concentrato sulle malattie genetiche con disregolazione immunitaria o autoinfiammazione e ho descritto come primo autore e autore corrispondente una nuova immunodeficienza primaria (JACI PMID 30771411). Ho condiviso come primo nome o autore corrispondente due pubblicazioni (JACI PMID 33556464 e 32437739) che rivelano la risposta al farmaco anti-IL-1R anakinra in pazienti con COVID19 (autorizzato dall'EMA).

Sono stato eletto per il periodo 2020-2024 come uno degli undici componenti (tesoriere) del comitato esecutivo della Società Europea di Immunodeficienza.

**Ho ottenuto come PI o collaboratore oltre 1,5 milioni di euro da finanziamenti competitivi**, permettendomi di finanziare la mia ricerca dal 2015. Ho supervisionato 5 post doc e 1 dottorando.

Sono stato invitato per presentazioni orali alla conferenza nazionale messicana di immunologia (2021), al meeting biennale della Pediatric Rheumatology European Society (2021), al meeting biennale della European Society of Immunodeficiency (2020-2022), alla conferenza nazionale della Società Italiana di Genetica Umana (2017) e il simposio annuale sull'autoimmunità, Harvard Medical School, Boston (2012).

## **Esperienze di lavoro**

### **Professore associato**

**Mar 2024- ad oggi**

Professore associato presso il dipartimento DINOEMI, Università degli Studi di Genova, Pediatra in convenzione presso l'Unità di Reumatologia e Malattie Autoinfiammatorie, Istituto Gaslini, Genova, Italia

### **Ricercatore a tempo determinato (RTDa,b)**

**Mar 2018- Feb 2024**

Ricercatore presso il dipartimento DINOEMI, Università degli Studi di Genova, Pediatra in convenzione presso l'Unità di Reumatologia e Malattie Autoinfiammatorie, Istituto Gaslini, Genova, Italia

### **Borsista di ricerca**

**Settembre 2014-Marzo 2018**

Ricercatore presso l'Istituto Gaslini, Genova, Italia, nel gruppo del Dott. Gattorno, che si occupa di immunodeficienze e interferonopatie di tipo 1 con responsabilità sia cliniche che di ricerca.

### **Collaboratore di ricerca**

**Settembre 2014-Agosto 2015**

Collaborateur de Recherche (studente post dottorato) presso l'Ospedale Universitario di Losanna, Losanna, Svizzera, nel laboratorio del Prof. Candotti: allestimento completo di un nuovo laboratorio, supervisione di uno studente di dottorato e sviluppo di un nuovo progetto sulla diagnosi delle interferonopatie di tipo I. Studio della disregolazione immunitaria in un modello murino della sindrome di Wiskott-Aldrich.

### **Research fellow**

**Aprile 2011 - Luglio 2014**

Borsista di ricerca post-dottorato presso il Children's Hospital di Boston, Harvard Medical School, nel laboratorio di immunologia pediatrica del Prof. Notarangelo, concentrandosi sullo studio delle immunodeficienze primarie attraverso un modello murino della sindrome di Wiskott Aldrich, un modello di zebrafish di una nuova immunodeficienza dovuta alla mutazione del gene EXTL3 in collaborazione con il Prof. Zon del Boston Children's Hospital e la generazione di linee cellulari iPS da pazienti con suscettibilità genetica all'encefalite da Herpes Simplex, in collaborazione con il Prof. Casanova della Rockefeller University, Len Zon del Boston Children's Hospital e la generazione di linee cellulari iPS da pazienti con suscettibilità genetica all'encefalite da Herpes Simplex, in collaborazione con il Prof. Casanova della Rockefeller University.

### **Borsista di ricerca**

**Settembre 2007- Aprile 2008**

Borsista presso l'Istituto di Ricerca in Biomedicina, Bellinzona, Svizzera, sotto la supervisione del Prof. Fabio Grassi, studio di un modello murino di lupus eritematoso sistemico.

### **Stagista di ricerca**

**Settembre 2006 - Agosto 2007**

Stagista di ricerca presso il CEBR, Centro di Eccellenza per la Ricerca Biomedica, Genova, sotto la supervisione della Dott.ssa Elisabetta Traggiai. Studio dell'effetto delle cellule staminali mesenchimali nelle malattie autoimmuni

## **Tirocinio medico**

**Agosto 2006**

Tirocinio presso l'Ospedale Universitario di Korle Bu, Accra, Ghana, presso il Dipartimento Pediatrico, sotto la supervisione del Dott. Onike Rodriguez.

## **Tirocinio di ricerca**

**Aprile 2004-Marzo 2005**

Stagista di ricerca presso il Laboratorio di Oncologia Sperimentale dell'Università di Utrecht, Paesi Bassi, e il Laboratorio Hubrecht per la biologia dello sviluppo, sotto la supervisione della Prof.ssa Rachel Giles e del Prof. Emile Voest, lavorando su un modello di angiogenesi per zebrafish.

## **Educazione e certificati clinici**

**Knowledge of Good Clinical Practice (GCP) online course** 13/03/2024

### **Esperienze in trial clinici:**

IVIG in hypogammaglobulinemia (site PI)

**Mar 2021**

### **Post Doctoral Student**

**Sett 2014 – Ago 2015**

Lausanne University, Lausanne University Hospital, Switzerland.

### **Post Doctoral Student**

**Gen 2011 – Lug 2013**

Harvard Medical School, Boston Children's Hospital, USA

### **PhD**

**Gen 2013 – Dic 2015**

PhD in Genetica, Università degli Studi di Genova. Tesi: Identification and characterization of a novel primary immunodeficiencies: *EXTL3* gene mutation causes a novel form of Immuno-Osseus dysplasia and unveils a critical role of heparan sulfate in thymopoiesis.

### **Specializzazione in pediatria**

**2007 - 2012**

Specializzato *cum laude* in Pediatria, Gaslini Children's Hospital, Genoa University, Italy.

### **Laurea**

**1999-2006**

Laura in Medicina e Chirurgia *cum laude*, con medaglia, Genoa University Medical School, Italy.

## **Premi**

### **Bright spark in immunology**

**September 2015**

European Federation of Immunological Societies, European conference of Immunology 2015.

## **Grants**

Fondo Italiano per la Scienza **Dec 2023-2028**

Combining gene editing and iPS to study and treat monogenic autoinflammatory diseases FIS00002235 953378 euro.

5 per mille Istituto Giannina Gaslini **June 2022-2025**

GASLINI genome-editing lab: development of an advanced cellular platform for the study of genetic diseases, 230000 euro.

Curiosity Driven Research Grant, Genova University **February 2021 – January 2023**

Combining CRISPR/Cas9 gene editing and induced pluripotent stem cells (iPS) to study and treat genetic diseases of the immune system, 65000 euro

Italian Health Ministry “Ricerca Finalizzata” (n. GR-2019-12369050) **February 2021 – January 2024**

Innovative pre-transplant conditionings to preserve bone marrow niche and improve the immunological reconstitution in immune dysregulation disorders, 80000 euro.

PRIN (n.20175XHBPN) Italian Research and University Ministry **Agosto 2019 – Agosto 2022**

Advanced genetic engineering to study and treat monogenic diseases, 140000 euro.

Italian Foreign Affairs Ministry (n. GR-2019-12369050) **February 2017 – January 2020**

Genetic basis of early-onset systemic lupus erythematosus in India and Italy, 146000 euro.

## **Presentazioni orali ed organizzazioni di congressi**

### **Oral presentation**

**April 2025**

International congress of Familial Mediterranean Fever and Systemic Autoinflammatory Disease, ISSAID, (Paris, France).

Systemic inflammation, lymphoproliferation and vasculopathy in a patient with ARHGAP10 mutation

### **Invited speaker**

**June 2023**

Inborn error working party symposium of the European Society for Bone Marrow transplantation (Brescia, Italy).

HSCT in ARPC1B and actinopathies

### **Invited speaker**

**October 2023**

European Society for Immunodeficiencies, biannual conference (Goteborg, Sweden)

Extinguishing the Fire: Inflammation Gone Wrong

### **Invited speaker**

**October 2021**

European Society for Immunodeficiencies, biannual conference (online)

ARPC1B related diseases

### **Invited speaker**

**February 2020**

Preceptorship on Autoinflammatory syndromes (Genova, Italy)  
Interferonopathies

**Oral presentation**

**June 2019**

International congress of Familial Mediterranean Fever and Systemic Autoinflammatory Disease, ISSAID, (Genova, Italy)

A combined immunodeficiency with severe infections, inflammation and allergy caused by ARPC1B deficiency

**Invited speaker**

**November 2019**

Maternal and fetal infections (Genova, Italy)

The immunology of pregnancy, of the fetus and of the newborn

**Invited speaker**

**October 2018**

Lupus 2018 (Florence, Italy)

Forme monogeniche di SLE.

**Oral presentation**

**June 2018**

Federation of Clinical Immunology Society (FOCIS) meeting (San Francisco, USA)

A combined immunodeficiency with severe infections, inflammation and allergy caused by ARPC1B deficiency

**Invited speaker**

**November 2017**

National conference of the Italian Society of Human Genetics (SIGU) (Napoli, Italy)

Genetic mechanisms in inflammatory diseases

**Invited speaker**

**October 2017**

La sindrome di Aicardi-Goutieres: un modello di interferonopatia (Brescia, Italy)

Interferonopatie di tipo 1 in reumatologia pediatrica: l'approccio diagnostico.

**Oral presentation**

**September 2017**

Pediatric Rheumatology European Society (PRES) annual meeting (Genova, Italy)

A severe case of polyarticular arthritis caused by mutation of the COPA gene.

**Invited speaker**

**May 2017**

Preceptorship on autoinflammatory syndromes (Genova, Italy)

Immunodeficiencies

**Eposter presentation**

**March 2017**

Clinical immunology society meeting (CIS) (Seattle, USA)

Efficacy of the JAK inhibitor ruxolitinib in two patients with SAVI syndrome

**Oral presentation**

**Sept 2016**

Pediatric Rheumatology European Society (PRES) meeting (Genova, Italy)

Type 1 interferonopathies: diagnostic role of peripheral blood interferon signature and preliminary results of treatment with a JAK 1/2 inhibitor.

**Oral presentation**

**April 2016**

Clinical Immunology Society (CIS) (Boston USA)

1) Blood Interferon signature as a screening for Type I Interferonopathies in children with early-onset SLE and vasculopathy.

2) Exostosin-like glycosyl transferase 3 (EXTL3) gene mutation causes a novel form of immuno-osseous dysplasia and unveils a critical role of heparan sulfate in thymopoiesis

**Oral presentation**

**October 2015**

International Congress of Familial Mediterranean Fever and Systemic Autoinflammatory Diseases, ISSAID (Dresden, Germany)

“Identification of type I interferonopathies using blood interferon signature: the experience of a pediatric rheumatology center”.

**Oral presentation**

**September 2015**

EUtrain-Translational Research in Rheumatology Conference (Genova, Italy)

“Identification of type I interferonopathies using blood interferon signature: the experience of a pediatric rheumatology center”.

**Oral presentation**

**Sept 2015**

Research day of Lausanne University Hospital (Lausanne, Switzerland)

“Selective deficiency of WASP in Treg cells is sufficient to cause autoimmunity in mice”.

**Oral presentations**

**Sept 2015**

European Conference of Immunology (Wien, Austria)

1) “Exostosin-like glycosyl transferase 3 (EXTL3) gene mutation causes a novel form of immuno-osseous dysplasia and unveils a critical role of heparan sulfate in thymopoiesis”

2) Selective deficiency of WASP in Treg cells is sufficient to cause autoimmunity in mice.

**Oral presentation**

**June 2015**

Annual European Congress of Rheumatology (EULAR), (Rome, Italy)

“Blood Interferon signature as a screening for Type I Interferonopathies in children with early-onset SLE and vasculopathy”.

**Oral presentation**

**May 2014**

Annual meeting of the Italian Paediatric Haematology and Oncology association (Genova, Italy)

“Role of Wasp and N-Wasp in b cell maturation, homing and function”

**Invited speaker**

**Oct 2012**

Annual symposium on Autoimmunity, Beth Israel Medical Centre, Harvard Medical School (Boston, USA) “*The autoimmunity of the Wiskott-Aldrich syndrome*”

### **Oral presentation**

**May 2012**

PRES course “Translational and Clinical issues in pediatric rheumatology” (Genova, Italy)

- B cell intrinsic deficiency of the Wiskott-Adrich syndrome protein (WASP) causes severe abnormalities of the peripheral B cell compartment in mice
- T cell activation dependent purinergic signalling in the pathogenesis of experimental lupus glomerulonephritis

### **Scientific meeting organization**

**Mar 2018**

Organizing committee for the 2019 meeting of the International Society of Systemic Autoinflammatory Diseases (ISSAID) (Genoa, Italy).

### **Scientific meeting organization**

**Dec 2010**

Scientific committee member of the Meeting on mitochondrial diseases (Gaslini Institute, Genoa, Italy).

## **Scientific association membership and reviewer activity**

Member of the European Society for Immune Deficiencies (ESID)

Review Editor for the journal “Frontiers in Medicine” and “Frontiers in Immunology”.

Reviewer for “Frontiers in Immunology”, “Journal of Allergy and Clinical Immunology”, “Clinical Immunology”, “Journal of Clinical Immunology”, “Clinical Cytometry”, “Current Medicinal Chemistry”, “Pediatric Rheumatology”.

Ministero della ricerca francese.

Accreditation for animal experimentation “Felasa EU functions ACD”.

## **Publications**

1. Woodward BL, Lahiri S, Chauhan AS, Garcia MR, Goodley LE, Clarke TL, et al. Inherited deficiency of DIAPH1 identifies a DNA double strand break repair pathway regulated by  $\gamma$ -actin. *Nature communications*. 2025;16(1).
2. Tesser A, Bocca P, Ulivi M, Pin A, Pastorino C, Cangelosi D, et al. Type I interferon signature: a quantitative standardized method for clinical application. *Clinical and Experimental Immunology*. 2025;219(1).
3. Simchoni N, Koide S, Likhite M, Kuchitsu Y, Kadirvel S, Law CS, et al. The common HAQ STING allele prevents clinical penetrance of COPA syndrome. *The Journal of experimental medicine*. 2025;222(4).

4. Palmeri S, Ferro J, Natoli V, Matucci-Cerinic C, Papa R, Rosina S, et al. Efficacy of High-Dose Intravenous Anakinra in Pediatric TAFRO Syndrome: Report of Two Cases and Literature Review. *Pediatric Blood and Cancer*. 2025.
5. Natoli V, Palmeri S, Rebollo-Giménez AI, Matucci-Cerinic C, Bocca P, Caorsi R, et al. Successful treatment of an anti-MDA5 antibody-positive Juvenile Dermatomyositis patient with refractory interstitial lung disease using tofacitinib. *Pediatric Rheumatology*. 2025;23(1).
6. Hadjadj J, Wolfers A, Borisov O, Hazard D, Leahy R, Jeanpierre M, et al. Clinical manifestations, disease penetrance, and treatment in individuals with SOCS1 insufficiency: a registry-based and population-based study. *The Lancet Rheumatology*. 2025;7(6):e391-e402.
7. Drago E, Fioredda F, Penco F, Prigione I, Bertoni A, Del Zotto G, et al. Inborn Error of WAS Presenting with SARS-CoV-2-Related Multisystem Inflammatory Syndrome in Children. *Journal of clinical immunology*. 2025;45(1).
8. Drago E, Bertoni A, Grossi A, Damasio MB, Anfigeno L, Miano M, et al. Majeed syndrome: first description in a patient of central-European ancestry. *Rheumatology*. 2025;64(5):3069-73.
9. Chan RWY, Serpas L, Ni M, Volpi S, Hiraki LT, Tam LS, et al. Erratum: Plasma DNA Profile Associated with DNASE1L3 Gene Mutations: Clinical Observations, Relationships to Nuclease Substrate Preference, and In Vivo Correction (The American Journal of Human Genetics (2020) 107(5) (882–894), (S0002929720303268), (10.1016/j.ajhg.2020.09.006)). *American Journal of Human Genetics*. 2025;112(5):1247.
10. Caorsi R, Bertoni A, Matucci-Cerinic C, Natoli V, Palmeri S, Rosina S, et al. Long-term efficacy of MAS825, a bispecific anti-IL1 $\beta$  and IL-18 monoclonal antibody, in two patients with systemic JIA and recurrent episodes of macrophage activation syndrome. *Rheumatology*. 2025;64(3):1528-33.
11. Al-Mayouf SM, Hadeef D, Aljaberi N, Movahedi N, AlEed A, Almutairi A, et al. A proposed clinical tool to identify high-risk patients for monogenic lupus: a pilot study. *Clinical and Experimental Rheumatology*. 2025;43(3):538-44.
12. Volpi S, Angelotti ML, Palazzini G, Antonelli G, Ravaglia F, Garibotto F, et al. Lupus Nephritis Patterns and Response to Type I Interferon in Patients With DNASE1L3 Variants: Report of Three Cases. *American Journal of Kidney Diseases*. 2024;84(6):791-7.
13. Papa R, Caorsi R, Volpi S, Gattorno M. Expert Perspective: Diagnostic Approach to the Autoinflammatory Diseases. *Arthritis and Rheumatology*. 2024;76(2):166-77.
14. Palmeri S, Penco F, Bertoni A, Bustaffa M, Matucci-Cerinic C, Papa R, et al. Pyrin Inflammasome Activation Defines Colchicine-Responsive SURF Patients from FMF and Other Recurrent Fevers. *Journal of clinical immunology*. 2024;44(2).
15. Orsi SM, Pepino C, Rossoni L, Serafino M, Caorsi R, Volpi S, et al. Corrigendum: Case report: Multisystem inflammatory syndrome in children with associated proximal tubular injury (Front. Nephrol., (2023), 3, (1194989), 10.3389/fneph.2023.1194989). *Frontiers in Nephrology*. 2024;4.
16. Naviglio S, Cicalese MP, Rivers E, Ferrua F, Bonfim C, Cenciarelli S, et al. Interleukin-1 blockade in patients with Wiskott-Aldrich syndrome: a retrospective multinational case series. *Blood*. 2024;144(16):1699-704.
17. McDonnell J, Cousins K, Younger MEM, Lane A, Abolhassani H, Abraham RS, et al. COVID-19 Vaccination in Patients with Inborn Errors of Immunity Reduces Hospitalization and Critical Care Needs Related to COVID-19: a USIDNET Report. *Journal of clinical immunology*. 2024;44(4).
18. Matucci-Cerinic C, Herzum A, Ciccarese G, Rosina S, Caorsi R, Gattorno M, et al. Therapeutic Role of HPV Vaccination on Benign HPV-induced Epithelial Proliferations in Immunocompetent and Immunocompromised Patients: Case Study and Review of the Literature. *Open Forum Infectious Diseases*. 2024;11(7).
19. Matucci-Cerinic C, Corona F, Varnier GC, Pastore S, Bocca P, Palmeri S, et al. Baricitinib treatment in children with COPA syndrome. *Journal of Allergy and Clinical Immunology: In Practice*. 2024;12(8):2201-4.
20. Hou C, Theodoropoulou K, Zaffalon L, Wang Z, Bertoni A, Volpi S, et al. HSP90 $\beta$  controls NLRP3 autoactivation. *Science Advances*. 2024;10(9).
21. Federici S, Cinicola BL, La Torre F, Castagnoli R, Lougaris V, Giardino G, et al. Vasculitis and vasculopathy associated with inborn errors of immunity: an overview. *Frontiers in Pediatrics*. 2024;11.
22. David C, Badonyi M, Kechiche R, Insalaco A, Zecca M, De Benedetti F, et al. Interface Gain-of-Function Mutations in TLR7 Cause Systemic and Neuro-inflammatory Disease. *Journal of clinical immunology*. 2024;44(2).
23. Coppola E, Sgrulletti M, Cortesi M, Romano R, Cirillo E, Giardino G, et al. The Inborn Errors of Immunity—Virtual Consultation System Platform in Service for the Italian Primary Immunodeficiency Network: Results from the Validation Phase. *Journal of clinical immunology*. 2024;44(2).
24. Chan YH, Lundberg V, Le Pen J, Yuan J, Lee D, Pinci F, et al. SARS-CoV-2 brainstem encephalitis in human inherited DBR1 deficiency. *Journal of Experimental Medicine*. 2024;221(9).

25. Cafaro A, Grossi A, Barco S, Pigliasco F, Biondi M, Schena F, et al. Diagnostic workflow for Adenosine Deaminase-2 Deficiency (DADA2): a proposal. *Biochimica Clinica*. 2024;48(1):62-6.
26. Cafaro A, Baiardi G, Pigliasco F, Barco S, Mattioli F, Volpi S, et al. A Novel LC-MS/MS Method for Therapeutic Drug Monitoring of Baricitinib in Plasma of Pediatric Patients. *Therapeutic Drug Monitoring*. 2024;46(1):67-72.
27. Caballero-Oteyza A, Crisponi L, Peng XP, Yauy K, Volpi S, Giardino S, et al. GenIA, the Genetic Immunology Advisor database for inborn errors of immunity. *Journal of Allergy and Clinical Immunology*. 2024;153(3):831-43.
28. Bruschi M, Angeletti A, Prunotto M, Meroni PL, Ghiggeri GM, Moroni G, et al. A critical view on autoantibodies in lupus nephritis: Concrete knowledge based on evidence. *Autoimmunity Reviews*. 2024;23(5).
29. Suter D, Bustaffa M, Papa R, Matucci-Cerinic C, Matarese S, D'Orsi C, et al. Corrigendum to "Clinical characterization, long-term follow-up, and response to treatment of patients with syndrome of undifferentiated recurrent fever (SURF)" *Seminars in Arthritis and Rheumatism* 55 (2022) 152024 (*Seminars in Arthritis and Rheumatism* (2022) 55, (S0049017222000750), (10.1016/j.semarthrit.2022.152024)). *Seminars in Arthritis and Rheumatism*. 2023;60.
30. Rossano M, Conti EA, Bocca P, Volpi S, Mastrangelo A, Cavalli R, et al. Novel heterozygous TREX1 mutation in a juvenile systemic lupus erythematosus patient with severe cutaneous involvement treated successfully with Jak-inhibitors: a case report. *Frontiers in Immunology*. 2023;14.
31. Orsi SM, Pepino C, Rossoni L, Serafino M, Caorsi R, Volpi S, et al. Case Report: Multisystem inflammatory syndrome in children with associated proximal tubular injury. *Frontiers in Nephrology*. 2023;3.
32. Matucci-Cerinic C, Malattia C, Pistorio A, Rosina S, Consolaro A, Viola S, et al. Skin manifestations help identifying different phenotypes of paediatric SAPHO syndrome. *Seminars in Arthritis and Rheumatism*. 2023;63.
33. Lancieri M, Bustaffa M, Palmeri S, Prigione I, Penco F, Papa R, et al. An Update on Familial Mediterranean Fever. *International Journal of Molecular Sciences*. 2023;24(11).
34. Hirschenberger M, Lepelley A, Rupp U, Klute S, Hunszinger V, Koepke L, et al. ARF1 prevents aberrant type I interferon induction by regulating STING activation and recycling. *Nature communications*. 2023;14(1).
35. Giardino G, Romano R, Lougaris V, Castagnoli R, Cillo F, Leonardi L, et al. Immune tolerance breakdown in inborn errors of immunity: Paving the way to novel therapeutic approaches. *Clinical Immunology*. 2023;251.
36. Fava D, Morandi F, Prigione I, Angelelli A, Bocca P, Pistorio A, et al. Blood Lymphocyte Subsets and Proinflammatory Cytokine Profile in ROHHAD(NET) and non-ROHHAD(NET) Obese Individuals. *Journal of the Endocrine Society*. 2023;7(9).
37. Dell'Orso G, Bagnasco F, Giardino S, Pierri F, Ferrando G, Di Martino D, et al. Hematopoietic stem cell transplantation for inborn errors of immunity: 30-year single-center experience. *Frontiers in Immunology*. 2023;14.
38. Benamar M, Chen Q, Chou J, Julé AM, Boudra R, Contini P, et al. The Notch1/CD22 signaling axis disrupts Treg function in SARS-CoV-2-associated multisystem inflammatory syndrome in children. *Journal of Clinical Investigation*. 2023;133(1).
39. Aldera E, Dufour F, Mercuri C, Rosina S, Volpi S, Gattorno M, et al. DADA-ism (2). *Medico e Bambino*. 2023;42(9):600.
40. Zhou Q, Kang G, Jiang P, Qiao R, Lam WKJ, Yu SCY, et al. Epigenetic analysis of cell-free DNA by fragmentomic profiling. *Proceedings of the National Academy of Sciences of the United States of America*. 2022;119(44).
41. Yang L, Booth C, Speckmann C, Seidel MG, Worth AJJ, Kindle G, et al. Phenotype, genotype, treatment, and survival outcomes in patients with X-linked inhibitor of apoptosis deficiency. *Journal of Allergy and Clinical Immunology*. 2022;150(2):456-66.
42. Wisniewski M, Chun A, Volpi S, Muscal E, Sexson Tejtzel SK, Munoz F, et al. Outcomes after SARS-CoV-2 Vaccination among Children with a History of Multisystem Inflammatory Syndrome. *JAMA Network Open*. 2022.
43. Trivioli G, Gelain E, Angelotti ML, Ravaglia F, Allinovi M, Lodi L, et al. A Report of 2 Cases of Kidney Involvement in ADA2 Deficiency: Different Disease Phenotypes and the Tissue Response to Type I Interferon. *American Journal of Kidney Diseases*. 2022;80(5):677-82.
44. Suter D, Bustaffa M, Papa R, Matucci-Cerinic C, Matarese S, D'Orsi C, et al. Clinical characterization, long-term follow-up, and response to treatment of patients with syndrome of undifferentiated recurrent fever (SURF). *Seminars in Arthritis and Rheumatism*. 2022;55.
45. Steiner A, Hrovat-Schaale K, Prigione I, Yu CH, Laohamonthonkul P, Harapas CR, et al. Deficiency in coatomer complex I causes aberrant activation of STING signalling. *Nature communications*. 2022;13(1).
46. Sin STK, Deng J, Ji L, Yukawa M, Chan RWY, Volpi S, et al. Effects of nucleases on cell-free extrachromosomal circular DNA. *JCI Insight*. 2022;7(8).

47. Signa S, Bertoni A, Penco F, Caorsi R, Cafaro A, Cangemi G, et al. Adenosine Deaminase 2 Deficiency (DADA2): A Crosstalk Between Innate and Adaptive Immunity. *Frontiers in Immunology*. 2022;13.
48. Penco F, Petretto A, Lavarello C, Papa R, Bertoni A, Omenetti A, et al. Proteomic Signatures of Monocytes in Hereditary Recurrent Fevers. *Frontiers in Immunology*. 2022;13.
49. Papa R, Rusmini M, Volpi S, Dell'Orso G, Giarratana MC, Caorsi R, et al. Progression of non-hematologic manifestations in SAMD9L-associated autoinflammatory disease (SAAD) after hematopoietic stem cell transplantation. *Pediatric Allergy and Immunology*. 2022;33(1).
50. Papa R, Caorsi R, Volpi S, Gattorno M. New monogenic autoinflammatory diseases: 2021 year in review. *Immunology Letters*. 2022;248:96-8.
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Autorizzo il trattamento dei miei dati personali in accordo con la legge GDPR 679/16 - "European regulation on the protection of personal data".

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