

**SINTESI CURRICULUM VITAE AI FINI DELLA PUBBLICAZIONE NELLA
SEZIONE AMMINISTRAZIONE TRASPARENTE**

FORMATO EUROPEO PER IL CURRICULUM VITAE


**INFORMAZIONI
PERSONALI**

Nome e Cognome	Giuseppe Agapito
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ESPERIENZA PROFESSIONALE	
Posizione Universitaria	
• Date (da – a)	Dal 02/03/2023 – Presente
• Nome e tipo di istituto di istruzione o formazione	Dipartimento di Giurisprudenza, Economia e Sociologia – Università degli Studi "Magna Græcia" di Catanzaro, Italia
• Qualifica conseguita	Professore Associato (SSD ING-INF/05)
Posizione Universitaria	
• Date (da – a)	2020 – 2023
• Nome e tipo di istituto di istruzione o formazione	Dipartimento di Giurisprudenza, Economia e Sociologia – Università degli Studi "Magna Græcia" di Catanzaro, Italia
• Qualifica conseguita	Ricercatore a Tempo Determinato (SSD ING-INF/05)

Istruzione e Formazione

Titoli di Studio Universitari	
• Date (da – a)	14/03/2025 a oggi
• Nome e tipo di istituto di istruzione o formazione	Ministero dell'Università e della Ricerca (MIUR), Italia
• Qualifica conseguita	Abilitazione Scientifica Nazionale alle funzioni di Professore Universitario di I fascia – Settore Concorsuale 09/H1 – Sistemi di elaborazione delle informazioni
Titoli di Studio e attività di aggiornamento post-universitari	
• Date (da – a)	2020
• Nome e tipo di istituto di istruzione o formazione	Ministero dell'Università e della Ricerca (MIUR), Italia
• Qualifica conseguita	Abilitazione Scientifica Nazionale alle funzioni di professore universitario di II fascia Settore concorsuale: ING-INF/05 – Sistemi di elaborazione delle informazioni Ministero dell'Università e della Ricerca (MIUR), Italia
Titoli di Studio e attività di aggiornamento post-universitari	
• Date (da – a)	11/09/2013
• Nome e tipo di istituto di istruzione o formazione	Università degli Studi "Magna Græcia" di Catanzaro, Italia Livello nella classificazione nazionale: Terzo ciclo / ISCED 8
• Qualifica conseguita	Dottorato di Ricerca in Ingegneria Biomedica e Informatica
Titoli di Studio e attività di aggiornamento post-universitari	
• Date (da – a)	2007
• Nome e tipo di istituto di istruzione o formazione	Università degli Studi della Calabria, Rende (CS), Italia

• Qualifica conseguita	Abilitazione alla professione di Ingegnere
Titoli di Studio e attività di aggiornamento post-universitari	
• Date (da – a)	2006
• Nome e tipo di istituto di istruzione o formazione	Università degli Studi della Calabria, Rende (CS), Italia
• Qualifica conseguita	Laurea in Ingegneria Informatica (Vecchio Ordinamento) Orientamento: Elettronica-Telecomunicazioni
Titoli di Studio e attività di aggiornamento post-universitari	
• Date (da – a)	1997
• Nome e tipo di istituto di istruzione o formazione	Istituto Tecnico Industriale Statale “A. Monaco”, Cosenza, Italia
• Qualifica conseguita	Diploma di Perito Industriale Capotecnico – Specializzazione Informatica (Progetto ABACUS)

MADRELINGUA	Italiano
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ALTRE LINGUE

	INGLESE
• Capacità di lettura	eccellente
• Capacità di scrittura	eccellente
• Capacità di espressione orale	eccellente

EVENTUALI ULTERIORI ALTRI TITOLI O ESPERIENZE VALUTABILI	Inclusione nella classifica mondiale “Single Year Top Scientists” Ente curatore: Stanford University Descrizione: Inserito nella classifica annuale dei Top 2% Scientists a livello mondiale (edizione 2023, basata su dati Scopus) per l’anno di riferimento, nella categoria <i>Single Year Impact</i> . La classifica è stata pubblicata il 4 ottobre 2023. Area scientifica: Informatica e Bioinformatica
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