## Programme (draft) of the 13th Bologna's convention on Crystal Forms - CF@Bo n. 13 University of Bologna 7-9 September 2025

	SUNDAY 7 September	Aula Prodi Complesso di San Giovanni in Monte University of Bologna	Abstract
11:30-14:00	REGISTRATION		
	Chairing Dario Braga	Opening of the convention	
14:00-14:30	<b>Fabrizia Grepioni</b> Department of Chemistry G. Ciamician, University of Bologna, Italy	Crystal Forms @ Bo	01
14:30-15:00	Alfred Lee MSD Research Laboratories Rahway, New Jersey 07065, USA	Solid Form Discovery: Past, Present and Future.	02
15:00-15:30	Philippe Fernandes Associate Director at Johnson and Johnson	Key Insights from Computational Tools in Solid-State Pharmaceutical Development	03
15:30-16:00	Franziska Emmerling Federal Institute for Materials Research and Testing (BAM) Department Materials Chemistry, Berlin	Real-Time Monitoring and Temperature Control for Optimized Polymorph Engineering	04
16:00-16:30	Ivo B. Rietveld SMS Laboratory University of Rouen Normandy	Preparation and stabilisation of metastable polymorphs for the improvement of API bioavailability.	05
16:30-17:00	Coffee break		
	Chairing Teresa Duarte		
17:00-17:30	Gareth Williams Department of Chemistry, University College London, UK	Accessing new polymorphs via solid solutions	O6
17:30-18:00	Helen Wheatcroft APS Crystallisation Scientist, AstraZeneca,UK	Crystallisation and Particle Control of a Multi-component API Crystal Form: From Structure to Process	07
18:00-18:30	Elena Simone Department of Applied Science and Technology (DISAT) Politecnico di Torino, Torino, Italy	Can crystal engineers make food? A few examples of crystallization strategies for the design of food formulations	08
18:30-19:00	Gerard Coquerel Laboratoire Sciences et Méthodes Séparatives. University of Rouen, France	Interferences between Polymorphism and Solid Solutions	09
19:00	Aperif in the courtyard	Complesso di San Giovanni in Monte University of Bologna	

	MONDAY 8	Hotel Aemilia	
	Chairing Gazhala Sadiq		
8:30-9:00	Jon Steed	Crystals as Intellectual Property	010
	Department of Chemistry, Durham University, Durham, UK		
9:00-9:30	Susan Reutzel-Edens	Turning polymorph challenges into patent opportunities	011
	SuRE Pharma Consulting, LLC		
9:30-10:00	Vania André	Unlocking New Antibiotic Forms: Crystal Engineering and	012
	IMS Researcher at CQE-IMS/IST	Supramolecular Strategies for Polymorphs, Cocrystals, and Beyond	
10:00-10:30	Sarah (Sally) Price	Pharmaceutical Digital Design: Can we go from Chemical	013
	Department of Chemistry, University College London, UK	Structure through Crystal Polymorph to Conceptual Crystallization Process?	
10:30-11:00	Coffee break		
	Chairing t.b.a.		
11:00-11:30	Marcus A. Neumann	A conceptual framework for the crystallizability of organic	014
	CEO Avant-garde Materials Simulation Deutschland GmbH,	compounds	
	Merzhausen, Germany		
11:30-12:00	Doris Braun	Hybrid Approaches in Solid Form Design: Virtual Screening and	015
	AssProf. PrivDoz. Dr. Institute of Pharmacy,	Experimental Validation	
	Christian Doppler Laboratory for Advanced Crystal Engineering		
	Strategies in Drug Development, University of Innsbruck		
12:00-12:30	Joost van den Ende	Machine Learning within CSP: from one crystal energy	016
	Roche Pharma Research and Early Development, Therapeutic	landscape to another	
	Modalities, Basel, Switzerland		
12:30-13:00	Rajni Miglani Bhardwaj	An integrated approach combining experimental and	017
	Associate research Fellow, Pfizer	computational for solid form design and selection	
13:00-14:00	Lunch and poster session		
14:00-16:00	Poster session	POSTER	Р
16:00-16:30	Coffee break		
	Chairing Matteo Daldosso		
		GOLD sponsor presentations	S

	Chairing t.b.a.		
17:30-18:00	Enrico Modena	Solid state, Polymorphism and the pharmaceutical	018
	Associate Director Science and Technology Novartis		L
18:00-18:30	Delia A. Haynes	Co-crystals, salts and sublimation	019
	Stellenbosch University		L
18:30-19:00	Thomas Rades and Inês Martins	Amorphous forms of drugs: from preparation to	O20
	Department of Pharmacy Faculty of Health and Medical	polyAmorphism	1
	Sciences University of Copenhagen		
19:00-19:30	Amy Woods-Ryan	HEPES of conformational, multi-zwitterionic polymorphs	021
	Durham University (PhD student) and GSK (Investigator)		
	Free evening in Bologna		
	TUESDAY 9 September	Hotel Aemilia	
	Chairing Lucia Maini		
8:30-9:00	Anna Slater	Continuous flow chemistry as a tool for crystallisation of	022
	University of Liverpool, Department of Chemistry and	porous organic materials	1
	Materials Innovation Factory		
9:00-9:30	Edyta Pindelska	Mechanisms of Cocrystal Formation and Coformer Exchange in	023
	Department of Pharmaceutical Chemistry and Biomaterials,	Ethenzamide Systems – From In Situ Studies to Pharmaceutical	1
	Medical University of Warsaw, Poland	Applications	
9:30-10:00	Matteo Daldosso	API: Don't Forget the I. From the Right Molecule to the Right	024
	Chief Scientific and Innovation Officer at PolyCrystalLine	Particle	
10:00-10:30	Federica Lazzari	Challenges in crystallization scale-up of an API nanocrystalline	025
	Solid state and crystallization scientist at Chemessentia Srl	form	1
	(part of Chemo group)		
10:30-11:00	Coffee break		
	Chairing Simone d'Agostino		
11:00-11:30	Joop ter Horst	Optimizing complex multicomponent solid form discovery &	026
	Tiofarma, NL	crystallization process design	
11:30-12:00	Irene Bassanetti	The Critical Role of 3D Molecular and Biomolecular Structures	027
	Senior Scientist in Analytics and Early Formulation	in Innovative Drug Discovery	1
	Department, Preclinical R&D Chiesi Farmaceutici		
12:00-12:30	Martin Viertelhaus	Solubility – Easy Parameter with Hurdles in Determination and	028
	Principal Scientist BASF SE - Analytical and Material Science	Interpretation	

12:30-13:00	Matteo Lusi Dpt of Chemical Science & Bernal Institute, University of	Crystalline Solutions for Pharmaceutical Problems	O29
	Limerick, Ireland		
13:00-14:00	Lunch		
	Chairing t.b.a.		
14:00-14:30	Luc Aerts Head Solid State Development Sciences, UCB Pharma	Co-crystallisation as a versatile tool in pharmaceutical development	O30
14:30-15:00	<b>Pietro Sacchi</b> Research and Application Scientist – The Cambridge Crystallographic Data Centre. UK	Computational approaches for the prediction of particle properties of organic molecular materials.	031
15:00-15:30	Joe Lubach Distinguished Scientist at Genentech, Inc., department of Synthetic Molecule Pharmaceutics.	Insights into Pharmaceutical Drug Substance and Product Using Multinuclear Solid-State NMR Spectroscopy	032
15:30-16:00	Helen Blade Principal Scientist - Solid State Computational Pharmaceutics, AstraZeneca	Connecting API to product	033
16:00-16:30	Coffee break and Poster Prizes		
16:30-17:30	NICE event: Nature Inspired Crystal Engineering		
16:30-16:45	Alessia Bacchi University of Parma	A NICE PoEM: cocrystals from liquid ingredients	O34
16:45-17:00	Giuseppe Resnati Polytechnic of Milan	Pharmaceutical cocrystals via halogen bond	O35
17:00-17:15	Michele Remo Chierotti University of Torino	Advanced Solid-State NMR tools for Crystal Engineering: From Structure Elucidation to Phase Purity Assessment	O36
17:15-17:30	Pavel Zolotarev University of Milan	Study of pillared MOFs with Zn-paddlewheel state switching	037
17:30	Closing CF@Bo n.13		

For queries about the programme refer to professor Dario Braga (dario.braga@unibo.it)