

DAY 1 – 03.10.2016	DAY 2 – 04.10.2016
08.30 – 09.00 Registration : Morning Coffee and Networking	08.30 – 09.00 Registration : Morning Coffee and Networking
09.00 – 09.20 Welcome, Intro to Speakers	
09.20 – 10.30 Commissioning & Qualification of a pharmaceutical facility: A new approach as per latest EU GMP Annex 15-Part 1 By Mr. Paolo Curto' , DOC Managing Director	09.00 – 10.30 NEW EP Monograph for WFI Production : a New Challenge for Pharmaceutical Industry By Mr. Paolo Curto' , DOC Managing Director
10.30 – 10.45 Morning Break	10.30 – 10.45 Morning Break
10.45 – 12.45 Commissioning & Qualification of a pharmaceutical facility: A new approach as per latest EU GMP Annex 15-Part 2 By Mr. Paolo Curto' , DOC Managing Director	10.45 – 12.45 Only MASCO can: Turn-key approach in a pharmaceutical facility,.
12.45 – 14.00 Lunch Break	12.45 – 14.00 Lunch Break
14.00 – 15.30 Process Validation: Extractables & Leachables Qualification of In-process materials and finished forms By Mr. Antonio Legnani-DOC Process Validation Director	14.00 – 16.30 Polysan: VISIT OF THE FACTORY Case study in real time
15.30 -15.45 Afternoon Break and Networking	
15.45 – 17.00 Process Validation: Sterilizing Grade Filter Validation By Mr. Antonio Legnani-DOC Process Validation Director	16.30 – 17.30 Q & A and Networking
17.00 -17.30 Q & A, End of Session	END OF THE TRAINING PROGRAM