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Andrew Lennard is in the Global Regulatory Affairs CMC team at Amgen and is based in the UK. Within CMC Regulatory Affairs, he is part of the External Engagement and Advocacy team with responsibilities in advancing approaches to accelerate CMC in product development. Andrew has over 15 years' experience in CMC Regulatory Affairs, with a special interest in control strategy and using prior knowledge, in which he has participated at the EMA workshops Prior Knowledge, **CMC** acceleration on and on Breakthrough/PRIME. He is also an active member of EFPIA leading several initiatives relating to CMC acceleration, including 'Stability' for which Andrew is the EFPIA topic lead on the Expert Working Group for the ICH revision of the stability guidelines. Prior to Regulatory Affairs, Andrew was a Principal Scientist in drug discovery for large pharma and small biotech start-up companies and holds a PhD from the University of Cambridge (UK).

