

Eliminating Human Error from Your List of Manufacturing Deviations



Ginette Collazo, Ph.D.

By

Mar 23, 2021 7:00 am EDT



To err is human, but is **human error** a justified “cause” in GMP investigations? In pharmaceutical manufacturing, upwards of 80% of process deviations can be attributed to human error. While regulatory agencies make clear that all deviations, including those caused by human error, must be fully investigated – can we say that we are effectively doing that as an industry? On today's episode of *Voices in Validation* we speak with Ginette Collazo as we dig deeper into the identification and reduction of human error in GMP facilities.

Last week to register with BEST pricing for Computer Systems Validation & Software Assurance Week Virtual Event

Source URL: <http://www.ivtnetwork.com/article/eliminating-human-error-your-list-manufacturing-deviations-0>