



Indoco's API manufacturing facility at Patalganga completes US FDA inspection

The facility successfully cleared the inspection with zero observations

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Indoco Remedies Limited announced today that the United States Food and Drug Administration (USFDA) successfully completed inspection of its Active Pharmaceutical Ingredients (API) manufacturing facility at Patalganga.

The inspection concluded with zero form 483 observations, reflecting the company's commitment to the highest standards of quality, regulatory compliance and operational excellence.

"This successful USFDA inspection is a testament to the strong quality culture and compliance in every process and every product. We remain committed to strengthening our systems to deliver safe and efficacious medicines across the globe," said Aditi Panandikar, Managing Director, Indoco Remedies Limited.