

# Elite Pharmaceuticals Receives FDA Approval for Codeine and Acetaminophen Combo

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**NORTHVALE, NJ / ACCESSWIRE / September 12, 2019 /** Elite Pharmaceuticals, Inc. ("Elite" or the "Company") (OTCQB:ELTP) a specialty pharmaceutical company developing abuse-deterrent opioids and niche generic products, today announced that it received approval from the US Food and Drug Administration (FDA) for an Abbreviated New Drug Application (ANDA) for a generic version of Tylenol® with Codeine (acetaminophen and codeine phosphate) 300mg/15mg, 300mg/30mg, and 300mg/60mg tablets.

## About Elite Pharmaceuticals, Inc.

Elite Pharmaceuticals, Inc. is a specialty pharmaceutical company which is developing a pipeline of proprietary pharmacological abuse-deterrent opioid products as well as niche generic products. Elite specializes in oral sustained and controlled release drug products which have high barriers to entry. Elite owns generic products which have been licensed to TAGI Pharma, Glenmark Pharmaceuticals, Inc., USA., and Lannett Company, Inc. Elite currently has thirteen approved generic products, three generic products filed with the FDA, one approved generic product pending manufacturing site transfer, and an NDA filed for SequestOx™. Elite's pipeline products include abuse-deterrent opioids which utilize the Company's patented proprietary technology and a once-daily opioid. These products include sustained release oral formulations of opioids for the treatment of chronic pain. These formulations are intended to address two major limitations of existing oral opioids: the provision of consistent relief of baseline pain levels and deterrence of potential opioid abuse. Elite operates a GMP and DEA registered facility for research, development, and manufacturing located in Northvale, NJ. Learn more at [www.elitepharma.com](http://www.elitepharma.com).

This press release contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995. Including those related to the effects, if any, on future results, performance or other expectations that may have some correlation to the subject matter of this press release, readers are cautioned that such forward-looking statements involve risks and uncertainties including, without limitation, Elite's ability to obtain FDA approval of the transfers of the ANDAs or the timing of such approval process, delays, uncertainties, inability to obtain necessary ingredients and other factors not under the control of Elite, which may cause actual results, performance or achievements of Elite to be materially different from the results, performance or other expectations that may be implied by these forward-looking statements. These forward-looking statements may include statements regarding the expected timing of approval, if at all, of SequestOx™ by the FDA, and the actions the FDA require of Elite in order to obtain approval of the NDA. These forward-looking statements are not guarantees of future action or performance. These risks and other factors, including, without limitation, Elite's ability to obtain sufficient funding under the LPC Agreement or from other sources, the timing or results of pending and future clinical trials, regulatory reviews, and approvals by the Food and Drug Administration and other regulatory authorities and intellectual property protections and defenses, are discussed in Elite's filings with the Securities and Exchange Commission, including its reports on forms 10-K, 10-Q, and 8-K. Elite is under no obligation to update or alter its forward-looking statements, whether as a result of new information, future events or otherwise.

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