

🕒 27 September 2009

GSK provides update on regulatory filings for Zunrisa/Rezonic

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


currently communicating with those authorities and all study investigators.

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GSK has made this decision, after regulatory consultation, based on the company's assessment that significant further safety data would be required to support the registration of casopitant on a worldwide basis, which would take a considerable time to produce. Consequently, all on-going regulatory files for casopitant are being withdrawn.

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


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