Unnatural fusion of API2 and MALT1 proteins leads to deregulated transcription in MALT lymphoma.

Printed from <u>https://www.cancerquest.org/newsroom/2011/01/unnatural-fusion-api2-and-malt1-proteins-leads-deregulated-transcription-malt</u> on 05/11/2024

Normal, healthy cells use specific proteins called transcription factors to regulate the activit of their genes. Importantly this is done in a tightly controlled manner. When the transcription factor NF- κ B is functioning normally, it is capable of being turned off and on, and thus cellular function is retained. In many instances of cancer, however, NF- κ B has lost its "off switch," which leads to unregulated gene activity (transcription).

Scientists at the University of Michigan have discovered a key factor leading to NF_KB deregulation in mucosa-associated lymphoid tissue (MALT) lymphoma. The proteins API2 and MALT1 are fused together to form an unnatural newprotein. This fusion oncoprotein binds to NF-_KB-inducing kinase (NIK) and cleaves it, thus removing the regulatory region of the kinase and leading to constant activity of the NF-_KB pathway. Neither API2 nor MALT1 appear capable of cleaving NIK on their own. It is only the fusion protein that has this activity.

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