

I. JAK kinases 與 IL-3 受體 α 和 βc 次單位結合之探討

II. p38 訊息傳遞於細胞分化與生長抑制之角色探討

I. Determinants for the interactions between the cytoplasmic regions of the Interleukin-3 receptor subunits α and βc and JAK kinases

and JAK kinases

II. Roles of the p38 pathway in cell differentiation and growth inhibition

中文摘要

IL-3 為細胞激素家族的一員，調控著細胞的生長、增生、分化與凋亡。IL-3 受體上沒有酪胺酸激酶，必須藉由 JAK 蛋白激酶傳遞訊息。我們實驗室過去以免疫沉澱法證實 JAK1 會與 IL-3 受體 β 次單位結合，而 JAK2 則與 IL-3 受體 α 次單位結合。而 JAK1 和 JAK2 與 IL-3 受體的結合是不受細胞激素刺激影響的。JAK 蛋白上有七個同源的區域，分別命名 JH1 至 JH7。目前已知 JH1 具有酵素活性，而 JH2 調控 JAK 的活性，而 JH3 至 JH7 片段則與受體的結合有關。但 JAK1 和 JAK2 究竟以哪個區域與 IL-3 受體結合？我利用實驗室先前建構好的 GST-IL-3 受體融合質體及一系列的 JAK1 和 JAK2 缺失質體，以 GST-pull down 方法分析。實驗結果顯示：JAK1 的 JH3 至 JH7 片段與 IL-3 受體 β 次單位結合，與 α 次單位有微弱的結合。JAK2 的 JH7 或 JH6 至 JH7 區域缺失的情況下，皆不影響與 IL-3 受體 β 次單位結合的能力，與 α 次單位有微弱的結合。

中文摘要

p38 pathway 對於調控細胞增生、分化及細胞凋亡扮演重要的角色。本實驗室過去研究發現，Activin A 誘導 globin gene promoter 活性及抑制細胞生長的作用會被 p38 抑制劑 SB203580 所抑制。SB203580 是專一性抑制 p38 α 和 p38 β 。為了想更進一步了解 p38 α 或 p38 β 參與細胞激素調控細胞增生與分化的情形，首先我利用 RT-PCR 分析在 K562 及 Jurkat 細胞內四個 p38 isoforms 表現的情形。在 K562 及 Jurkat 細胞中 p38 α , p38 β , p38 γ 和 p38 δ 皆有表現。我們的實驗結果發現 Activin A 所誘導的 globin 和 globin gene promoters 活性會被 p38 dominant negative mutants p38 α AF 和 p38 β AF 所抑制，顯示 Activin A 誘導 \sim globin 和 \sim globin gene promoters 活性的作用是透過 p38 α 和 p38 β 。另外，c-Jun，紅血球分化的抑制者，其蛋白表現量於 K562 細胞中會受 SB203580 正向調控。最後，我利用 Interferon alpha (IFN- α) 刺激 Jurkat 細胞，發現 IFN- α 會透過 p38 pathway 抑制 Jurkat 細胞生長。綜合這些結果，於 K562 和 Jurkat 細胞中 p38 扮演調控細胞生長和分化的重要角色。

英文摘要

IL-3, a member of cytokine family, is involved in the cell growth, proliferation, differentiation and apoptosis. IL-3 stimulates its biological effects by binding to a heterodimeric receptor composed of α and β subunits. JAK1 and JAK2 are rapidly activated after binding of IL-3 to their receptors, and are essential for the initiation of intracellular signaling. Previously, we reported that JAK1 and JAK2 were associated with the IL-3 receptor β and α subunits, respectively. In addition to the JH1 domain (kinase domain), the JAK family contains the JH2-JH7 domains. JH2 has been shown to have a negative regulatory effect on JAK2 kinase activity. JH3-JH7 domains have been implicated in receptor association. However, which regions of JAKs interact with IL-3 receptors is now unclear. In this study, I demonstrated that the JH3-JH7 domains of JAK1 interacted with β c subunit, but had weak interaction with IL-3 receptor α subunit by GST-pull down assay. The deletion of JH7 domain or JH6-JH7 domains of JAK2 did not affect the interaction with IL-3 receptors.

Abstract

p38 play a central role in cellular responses such as cell proliferation, differentiation and apoptosis. We have previously reported that Activin A-induced globin promoters and cell growth inhibition were inhibited by p38 inhibitor SB203580. The p38 inhibitor SB203580 targets both p38 α and p38 β . To further investigate that p38 α , p38 β or both kinases are involved in cytokine-mediated cell proliferation and differentiation, we performed the RT-PCR to analyze the expression of four p38 isoforms on K562 and Jurkat cells. We showed that p38 α , p38 β , p38 γ and p38 δ transcripts are expressed in K562 and Jurkat cells. Expression of the exogenous p38 dominant negative mutants, p38 α AF and p38 β AF, reduced Activin A-induced α - and ζ -globin promoters activity, indicating that Activin A activated α - and ζ -globin gene promoters through p38 α and p38 β . In addition, the expression of c-Jun, a blocker of erythroid differentiation, was up-regulated by SB203580 in K562 cells. At last IFN- α inhibited Jurkat cell growth through p38 pathway. Together, these results suggest that p38 plays an important role in regulating cell proliferation and differentiation.