• 系統編號	RC8901-0140		
• 計畫中文名稱	環境中氦氧化合物與亞硝基化合物之毒理學研究氧化氦基因毒性分子機制之探討(III)		
• 計畫英文名稱	Studies on the Molecular Mechanisms of Genotoxicity of Nitric Oxide (NO) (III)		
• 主管機關	行政院國家科學委員會	• 計畫編號	NSC88-2621-B038-002-Z
• 執行機構	台北醫學院醫事技術系		
• 本期期間	8708 ~ 8807		
• 報告頁數	0 頁	• 使用語言	英文
• 研究人員	何元順;林仁混 Ho, Yuan-Soon;Lin, Jen-Kun		
• 中文關鍵字	基因毒性;一氧化氮;細胞凋亡;抗氧化劑;環境衛生		
• 英文關鍵字	Genotoxicity; Nitric oxide (NO); Apoptosis; Antioxidant; Environmental health		
• 中文摘要	NO 在以前一直被認為僅從燃燒的廢氣產生,是一種體外的汙染物,並不受研究者的重視。最近生物醫學的進步,藥理與生化學家研究出 NO 是一種重要的內生性傳遞物質,它可在自體細胞合成,具有很強的生物功能,諸如血管的擴張鬆弛,血壓的調控,神經訊息的傳遞,基因的表現等等。其重要性很受注目,1992 年 Science 雜誌就把 NO 選爲年度風雲分子(The Molecular of the Year)。自此這個不起眼的簡單分子,就搖身一變爲大名鼎鼎,如雷貫耳的奇異分子。GSNO(S-nitrosoglutathione)爲細胞內抗氧化物 GSH 攜帶一氧化氮之化合物,當它釋出一氧化氮時,可對細胞造成 DNA 傷害。已有許多報告指出一氧化氮自由基可造成細胞凋亡,但 GSNO 引起細胞凋亡之詳細機制尚未被研究清楚。在本實驗中,我們使用人類腸癌細胞株,來探討 GSNO 對細胞之毒性及細胞凋亡之機制。我們發現,細胞給予 GSNO 處理之同時,若一起加入銅離子(Cu/sup ++/)則細胞死亡率較單獨處理 GSNO 明顯減少。據此我們亦證實了 Cu/sup ++/在細胞外可促使 GSNO 釋出 NO ? ,由於 NO ?不穩定,在極短時間內轉變爲硝酸鹽及亞硝酸鹽,因而降低進入細胞內 NO ? 之量,而減少細胞的傷害。NO ? 調控細胞凋亡過程中之基因表現,是我們想要嘗試了解的,以西方墨點分析來偵測基因表現的變化,結果發現 Bad、Bax 及 c-Jun 等蛋白的表現皆有增加之現象,而 p27 及 Bcl-2的表現卻有被抑制的現象。爲了解 NO ? 在參與細胞內訊息傳遞所扮演之角色,我們也對 PKA 及 PKC 之表現進行探討,結果發現 PKA 與 PKC.zeta.之表現會被 NO ? 抑制,此結果告訴我們在 GSNO 引起之細胞凋亡中,存在一種特殊的調節機制,而 PKA 及 PKC.zeta.可能扮演一重要角色。		

• 英文摘要 In this study, the amount of S-nitrosoglutathione (GSNO) was measured spectrophotometrically at 334nm. Spontaneous decrease of

absorbency at 334nm was detected when GSNO was exposed to 37.degree.C and a high pH (pH 8.0). We investigated the catalytic roles of various metal ions on the decomposition of GSNO. The degradation of GSNO (0.5mM) was enhanced by the presence of Cu/sup 2+/ and Ni/sup 2+/ ions. The amount of NO release from GSNO degradation was estimated by the Griess reaction based on nitrite accumulation. The results indicate that nitrite production was elevated by at least 2-fold in the presence of Cu/sup 2+/. Our study further indicates that Cu/sup 2+/ enhance GSNO-induced apoptosis in human colon adenocarcinoma (HT 29) cells. We also found that copper ions modulate the expression of bad, bax, and bcl-2 in GSNO-treated HT 29 cells. The levels of bax and bad proteins were significantly elevated by about 4- to 6-fold when compared with mock-treated cells at 24 h after combined treatment of GSNO plus Cu/sup 2+/ or Ni/sup 2+/. On the other hand, significant inhibition of bcl-2 occurred in HT 29 cells with simultaneous treatment of GSNO with Cu/sup 2+/ (or Ni/sup 2+/). It seemed that Cu/sup 2+/ (Ni/sup 2+/) could enhance the decomposition of GSNO that liberated NO to activate the pathways. Our results demonstrated that the apoptotic effects induced by GSNO was promoted by Ni/sup 2+/ and Cu/sup 2+/ through two different mechanisms: by depletion of intracellular GSH level and by triggered of NO release from GSNO which then promoted the NO-induced apoptotic cell death in human cells.