Hypotension induced by interleukin-3 in patients on angiotensin-converting enzyme inhibitors

Sir—The use of haematopoietic growth factors is increasing. It is, therefore, essential to demonstrate their efficacy and safety when given with other commonly prescribed medicines. We treated 26 patients with ovarian or small-cell undifferentiated cancers with chemotherapy followed by recombinant human interleukin-3 (rhIL-3) (0·125–7·5 μg/kg daily).1 3 of these patients were also receiving angiotensin-converting enzyme (ACE) inhibitors for hypertension. All 3 developed marked hypotension (WHO toxicity grade 2 or 3) within 1–4 hours of the first rhIL-3 injection. In all 3 patients discontinuation of the ACE inhibitor while continuing rhIL-3 resulted in return of the blood pressure to normal. After stopping rhIL-3 the blood pressure increased and ACE inhibitors were again required. Hypotension was not observed in 22 patients given rhIL-3 without ACE inhibitors. 1 other patient, not receiving an ACE inhibitor, became hypotensive during a period of neutropenic fever. The fever resolved with antibiotic therapy.

The hypotensive effects of ACE inhibitors are mediated by inhibiting the conversion of angiotensin I into angiotensin II (a powerful vasoconstrictor). Additionally, inhibition of kinase II reduces the breakdown of bradykinin, resulting in vasodilation.2 Furthermore, rhIL-3 stimulates the synthesis of secondary cytokines (eg, IL-6)3 and histamine.4 The hypotension induced by these substances is mediated via nitric oxide (NO). Both bradykinin and histamine stimulate rapid (within minutes) NO production by activation of constitutive NO synthase. Cytokines, including IL-6, cause a late and more sustained NO generation via an inducible NO synthase found in endothelium and vascular smooth muscle.5

The urine from 1 of the hypotensive patients was assayed for nitrate and cGMP, as indicators of in-vivo NO production. In the first chemotherapy cycle without rhIL-3 the 24-hour urinary nitrate and cGMP excretion did not exceed 1 mmol and 1 μmol, respectively. In the second cycle with rhIL-3, 24-hour nitrate excretion rose from 0·9 mmol before chemotherapy and 0·8 mmol before rhIL-3 to 5·7 mmol on day 5 of rhIL-3 administration. cGMP rose from 0·9 μmol to 2·6 μmol on day 5. In another 16 patients, not receiving ACE inhibitors and not developing hypotension, 24-hour nitrate was measured in 9 and 26 cycles of chemotherapy with and without rhIL-3, respectively. In none of these cycles did the 24-hour nitrate excretion exceed 1·3 mmol (control cycles 0·7 [SE 0·3] mmol; cycles with rhIL-3 0·8 [0·3] mmol).

These data suggest that NO generated in vessel walls by bradykinin and histamine (via constitutive NO synthase) and secondary cytokines (via inducible NO synthase) were responsible for the increase of urinary nitrate and cGMP. Without priming by ACE inhibitors, the effects of rhIL-3 were too small to lower blood pressure or could easily be compensated by counter-regulatory mechanisms. We suggest that synergism between forces acting to generate NO, in the absence of the counteracting effects of angiotensin II, were responsible for the rapid induction of hypotension in patients treated with ACE inhibitors combined with rhIL-3.

European Cancer Centre, Amsterdam; *Department of Oncology, Free University Hospital, 1081 HV Amsterdam, the Netherlands; Department of Surgery, Free University Hospital; and Department of Medical Oncology, the Netherlands Cancer Institute/Antoni van Leeuwenhoekhuis, Amsterdam

Nitric oxide in ulcerative colitis

Sir—Lundberg and colleagues (Dec 17, p 1673) suggest that nitric oxide (NO) concentrations in the colonic lumen might be increased in ulcerative colitis. The chemiluminescence method used by these workers depends on the release of energy from the reaction of NO with ozone, and is therefore not truly direct. Ozone is a reactive species and the colonic lumen contains many gaseous substances, so there is doubt as to its specificity under these conditions. H₂S in particular may react with ozone to produce chemiluminescent species, and is known to be raised in ulcerative colitis.

We have evidence that the concentration of NO might be increased in ulcerative colitis, with a molecule-specific technique. The rectal lumen in 8 patients with active disease and 8 controls was perfused for 1 h with a nitrogen stream, and NO was collected in a cold trap at −196°C. The NO thus obtained was measured by infrared diode laser spectroscopy. This technique allows high detection sensitivity and specificity and we have checked that the following gases have no absorptions overlapping the NO transitions used in the measurements: H₂O, H₂S, CH₄, CO, CO₂, NO, N₂O. The estimated minimum detectable amount of NO was 0·1 nmoL. NO was detected in 4 of 8 patients with active ulcerative colitis but in none of the controls (p<0·05). The amounts measured varied from 0·13 to 1·1 nmoL. Patients negative for NO all had blood visible in the rectum and the collection tubing.

NO is avidly bound by haemoglobin and we suspect that this was why we could not detect it in 4 of our patients. If the technique used by Lundberg and co-workers were truly specific for NO, we would have expected them to encounter the same difficulty. Nevertheless, Lundberg and colleagues’ work supports our observation that NO synthesis is increased in ulcerative colitis,1 a finding that could prove to be of both pathogenetic and therapeutic importance.

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* P D Reynolds, S J Middleton, G M Hansford, J O Hunter
*Gastroenterology Research Unit, Addenbrooke’s Hospital, Cambridge CB2 2QZ, UK; and Department of Chemistry, University of Cambridge


Sir—Lundberg and colleagues’ study complements our own investigation showing raised serum concentrations of NO metabolites in 26 patients with severe attacks of ulcerative colitis.1 However, in that study concentrations of C-reactive protein were shown to predict clinical outcome more accurately than NO metabolites. Lundberg’s data concern an investigation showing raised serum concentrations of NO metabilites in 26 patients with severe attacks of ulcerative colitis.1 This was why we could not detect it in 4 of our patients. If the technique used by Lundberg and co-workers were truly specific for NO, we would have expected them to encounter the same difficulty. Nevertheless, Lundberg and colleagues’ work supports our observation that NO synthesis is increased in ulcerative colitis,1 a finding that could prove to be of both pathogenetic and therapeutic importance.

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