Gadolinium and systemic fibrosis: guilt by association

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Nephrogenic systemic fibrosis (NSF, originally called ‘nephrogenic fibrosing dermopathy’) is a painful and debilitating fibrosing disorder that was first identified in 1997 in several patients with stage 5 chronic kidney disease (glomerular filtration rate (GFR) <15 ml/min/1.73 m² or permanently requiring dialysis) who had undergone renal transplantation at Sharp Memorial Hospital in San Diego, California.1 It typically manifests with skin tightening, tethering and hyperpigmentation on the trunk and on the extremities, progressing from distal to proximal, and often results in fixed flexion contractures of the fingers, elbows, knees and ankles. More recently, extensive systemic involvement has been identified including fibrosis of lymph nodes, thyroid, oesophagus, heart, lungs, liver, diaphragm, skeletal muscle, genitourinary tract and dura mater.2–3 The presence of skin changes of NSF has been associated with a nearly threefold increased risk of death within 24 months.4

On skin biopsy, the dermis is hypercellular with increased numbers of dermal fibroblasts.5–6 Thickened collagen bundles aligned with intact elastic fibres extend from the superficial dermis through the subcutis along interlobular septa.5 Interstitial mucin deposition is increased. CD34 fibrocytes, presumably originating from the circulation, and occasional cells of monocyte–macrophage lineage that stain for factor XIIIa and CD68 may be present in the dermis. When deeper tissue is sampled, fibrosis of subcutaneous tissue and fascia can also be observed.7 However, unlike scleroderma or scleromyxoedema, there is little or no inflammatory infiltrate in the skin of patients with NSF. Although the skin changes and joint contractures of NSF may resemble those of chronic graft-versus-host disease, NSF occurs in the absence of recent allogeneic haemopoietic cell transplantation; on skin biopsy, increased interstitial mucin and CD34 fibrocytes in the dermis typically differentiate NSF from chronic graft-versus-host disease.8

In 2006, Grobner reported that five of nine patients receiving haemodialysis in his unit developed NSF within days to weeks after undergoing MRI with Omniscan, a gadolinium-containing contrast agent.9 At that time, because of the lower risk of nephrotoxicity, gadolinium-containing contrast agents were assumed to be safer than iodinated contrast agents for administration to patients with chronic kidney disease.10 Thus, gadolinium-enhanced MR angiography of the lower extremities was often performed in patients with stage 5 chronic kidney disease to evaluate the condition of the peripheral arteries prior to renal transplantation.11 Gadolinium-containing contrast agents are composed of the rare earth metal gadolinium non-covalently bound to an organic chelating agent, with excess chelate added to bind free gadolinium that might have dissociated from the gadolinium–chelate complex. Omniscan is an aqueous solution that contains gadodiamide and excess free chelate caldiamide sodium. Magnevist is an aqueous solution that contains gadopentetate dimeglumine and excess free chelate diethylene-triamine pentaacetate. Grobner hypothesised that metabolic acidosis might destabilise the gadolinium–chelate complex, resulting in dissociation of gadolinium from chelate and deposition of gadolinium in tissue.

Following Grobner’s insightful suggestion that NSF might be a complication of gadolinium-containing contrast agent administration in patients with stage 5 chronic kidney disease, several lines of investigation have established a strong causal relationship between exposure to gadolinium-containing contrast agents used in imaging procedures and the subsequent development of NSF. The prevalence of NSF among patients with stage 5 chronic kidney disease who had been exposed to a gadolinium-containing contrast agent has been estimated to be as high as 18%12 to 30%.4 In a prospective case–control cohort study, Todd and colleagues observed skin changes characteristic of NSF in 13% of 186 patients with stage 5 chronic kidney disease who were receiving haemodialysis in community-based centres. Among these patients, prior exposure to a gadolinium-containing contrast agent was strongly associated with the subsequent development of NSF skin changes (OR 14.7).4 The risk of developing NSF was both time- and dose-dependent in a different cohort of 36 patients with stage 5 chronic kidney disease and biopsy-proven NSF who had been exposed to Magnevist: NSF developed within 3 months after the last dose in 66% of patients and the likelihood was greater with higher cumulative and total doses of the gadolinium-containing contrast agent (OR 1.2).13 Gadolinium has been detected and quantified in biopsies of skin14 and other tissues3 from patients with NSF and in the skin of rats, which previously had been subjected to 5/6 nephrectomy, after the intravenous administration of a gadolinium-containing contrast agent.15 Thus, the association between administration of gadolinium-containing contrast agents to individuals with chronic kidney disease and the subsequent development of NSF satisfies most of the Bradford–Hill criteria for causality.16 Demonstration of a plausible mechanism by which a gadolinium-containing contrast agent could induce the pathological changes of NSF would further substantiate this cause and effect relationship.

A clue to the pathophysiological mechanism by which tissue fibrosis develops after administration of a gadolinium-containing contrast agent is derived from the observation that skin thickening and tethering improve dramatically when patients with NSF are treated with imatinib mesylate.17 This small molecule tyrosine kinase inhibitor blocks signalling through both the transforming growth factor β (TGFβ) receptor and the platelet-derived growth factor (PDGF) receptor in normal and scleroderma dermal fibroblasts in vitro, thereby decreasing transcription and translation of type I collagen and fibronectin genes.18 In a murine model of bleomycin-induced dermal fibrosis, pretreatment with imatinib mesylate reduces extracellular matrix protein synthesis and dermal thickness18 and treatment with imatinib mesylate, beginning 3 weeks after the

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first intracutaneous bleomycin injection, not only stops further progression but also induces regression of pre-existing dermal fibrosis. TGFβ expression is increased in affected skin and muscle from patients with NSF compared with healthy controls. Incubation of normal human peripheral blood monocytes with either Omniscan, gadopentetate dimeglumine or gadolinium chloride stimulates the expression of a number of cytokines and growth factors, including TGFβ. Thus, exposure of patients with chronic kidney disease to gadolinium-containing contrast agents may induce the production of TGFβ and possibly of PDGF.

In this issue of Annals of the Rheumatic Diseases, two original reports from the same group describe additional effects of gadolinium-containing contrast agents on fibroblast and macrophage function that further contribute to understanding how a rare earth metal chelated by an organic ligand may induce tissue fibrosis. Piera-Velazquez and colleagues report direct effects of Omniscan and gadopentetate dimeglumine on dermal fibroblasts that are relevant to the pathogenesis of NSF. They found a marked increase in type I collagen production after culture of normal dermal fibroblasts for 48 h with either gadolinium-containing contrast agent but not with the free chelate chelidamine sodium. They also showed markedly increased production of the extracellular matrix components types I and III collagen, fibronectin and hyaluronic acid by dermal fibroblasts cultured from patients with NSF compared with those from healthy controls. In three patients with NSF, fibroblasts cultured from clinically affected skin displayed an activated myofibroblast phenotype and produced greater amounts of COL1A1 mRNA and type I procollagen and collagen proteins than did fibroblasts cultured from clinically unaffected skin. After serial passages of these NSF dermal fibroblasts in culture, an ‘activated’ phenotype with increased expression of genes encoding extracellular matrix macromolecules was maintained. Taken together, these data suggest that persistent activation of dermal fibroblasts, perhaps resulting from a direct action of free gadolinium or of the gadolinium–chelate complex on the fibroblast, may account for the fibrosis observed in patients with NSF. These observations will be strengthened if the phenotypic changes reported here in dermal fibroblasts cultured from patients with NSF can be reproduced by exposure of ‘normal’ dermal fibroblasts to gadolinium-containing contrast agents in vitro.

Del Galdo and colleagues observed differential expression of 551 genes when normal human monocyte-derived macrophages exposed to Omniscan were compared with macrophages exposed to phosphate buffered saline. They define a ‘macrophage Gd signature’ as those 31 upregulated genes with the highest signal intensity, of which three are interferon-inducible genes and five encode CC and CXC chemokines, most prominently CCL8/MCP-2. They extended these observations to NSF by demonstrating markedly increased CCL8/MCP-2 expression in affected skin from three patients with NSF compared with that in normal skin obtained from a healthy control individual. Common to the production of these chemokines is activation of nuclear factor κB (NFκB). By inhibiting NFκB with IκK-NBD, a cell-permeable NFκB inhibitory peptide, they completely abrogated Omniscan-induced upregulation of CCL8, CXCL10 and CXCL11 and markedly reduced the upregulation of CCL12. These findings provide further evidence that a gadolinium-containing contrast agent can directly alter human monocyte-derived macrophage function in a way that could contribute to the development of NSF. Further investigation should be directed towards clarifying the mechanism by which gadolinium interacts with cells to trigger these changes in function, so that interventions might be developed to inhibit these processes and prevent the development of NSF following exposure to a gadolinium-containing contrast agent.

Since NSF was first observed over a decade ago, mounting evidence has associated the development of this devastatingly disabling condition with exposure of patients with underlying kidney disease to gadolinium-containing contrast agents used in imaging procedures. Thus, it is now time to rename this condition more precisely as ‘gadolinium-associated systemic fibrosis’ and to remove ‘nephrogenic’ from its name, since NSF neither originates in the kidney nor is caused by factors originating in the kidney. At present, no method is available to prevent gadolinium deposition in tissue; once it has deposited, gadolinium persists in tissue. Radiologists have developed guidelines to try to decrease the incidence of this condition, which include measuring the serum creatinine level before administering a gadolinium-containing contrast agent to an individual at risk for impaired renal function, avoiding the use of Omniscan or Magnevist in patients with GFR <30 ml/min/1.73 m² and injecting the lowest possible dose of a gadolinium-containing contrast agent when no alternative exists. However, the combined epidemiological and laboratory data convincingly support avoiding the use of these drugs in patients with renal impairment. Only by exercising tremendous restraint when deciding whether or not to administer a gadolinium-containing contrast agent to a patient with diminished renal function can this iatrogenic epidemic be brought to an end.

References
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