ABSTRACT: A 72-year-old woman presented with painful proptosis of the right eye and a large destructive tumour of the middle cranial fossa. A diagnosis of IgDK multiple myeloma was made, based on histopathologic and immunologic studies of the biopsy. Biochemistry and bone marrow examination further confirmed the myeloma as IgDK type. The clinical, radiological, and pathologic findings are presented. The patient was treated with radiotherapy with satisfactory results.

RÉSUMÉ: Myelome multiple IgDK se présentant par une proptose unilaterale Nous rapportons le cas d'une femme de 72 ans qui s'est présentée avec une proptose douloureuse de l'œil droit et une grosse tumeur destructrice de la fosse crânienne moyenne. Le diagnostic de myélome multiple type IgDK fut posé sur la base d'études histopathologiques et immunologiques de la biopsie. La biochimie et l'examen de la moelle osseuse ont également confirmé le type de myélome. Nous en présentons les aspects cliniques, radiologiques et pathologiques. La patiente fut traitée par radiothérapie avec des résultats satisfaisants.

Multiple myeloma is a plasma cell dyscrasia frequently associated with multiple lytic bone lesions, anemia, monoclonal immunoglobulin in the serum, and commonly monoclonal light chains in the urine. It is known that IgD multiple myeloma comprises about 2% of all cases of myeloma and that in 89% of these cases, the light chains are of lambda type (Pruzanski and Rather, 1970). Orbital involvement by myeloma occurs in 0.3% of cases (Hayes et al., 1980) and the majority are unilateral. We recently encountered a patient with IgD multiple myeloma presenting with unilateral painful proptosis.

CASE REPORT

A 72-year-old woman was admitted to Sunnybrook Medical Centre on March 6, 1984 with a two month history of painful proptosis of the right eye. She had complained of severe low back pain six months prior to admission, at which time x-rays of the lumbosacral spine revealed diffuse bone thinning with compression of the L1 vertebral body. A provisional diagnosis of osteoporosis was made, and she was treated with acetaminophen, codeine, calcium supplement, and Naprosyn. Over the next six months, she developed anorexia, nausea and vomiting, and lost 13.6 kg.

Examination on admission revealed a thin elderly woman with a slightly tender, soft tissue mass above the right zygoma. There was right-sided exophthalmos (Hertel exophthalmometer reading of 21 mm. on the right and 12 mm. on the left). Corrected visual acuity on the right was 20/100, and on the left 20/50. Adduction and upward gaze of the right eye was limited. The optic fundi were normal, and the corneal reflexes were intact. The remainder of the neurological examination was normal. There was tenderness of the upper lumbar vertebrae. Otherwise, the general physical examination was normal.

Orbital x-ray showed destruction of the greater and lesser wings of the right sphenoid bone. Orbital ultrasound demonstrated a 1.7 cm. solid mass posterior to the orbit, lateral to the lateral rectus muscle and displacing it medially. Contrast-enhanced CT scan showed a large uniformly enhancing tumour on the right side destroying the floor of the middle cranial fossa, and extending upward to displace the temporal lobe. There was also destruction of the lesser wing of the sphenoid, postero-lateral orbital wall, and a portion of the lateral orbital roof. Inferiorly, the tumour extended along the pterygoid muscles and destroyed the pterygoid plates, with extension into the nasal cavity, nasopharynx, and sphenoid sinus (Figure 1). Skeletal survey showed compression of L1 vertebral body and loss of posterior elements of L5 (Figure 2). There were also multiple lytic bone lesions of the skull, both hemiuri, both femurs, and ribs.

A biopsy was taken from the right temporal bone where it was eroded by grayish tumour. The histologic and electron microscopic appearance was typical of multiple myeloma. Immunoperoxidase staining revealed that it was of the IgDK type (Figure 3).

The hemoglobin was 11.0 g/l with normal leukocytes and platelets. The BUN was 5 mmol/l (N = 3.0 - 7.0 mmol/l), creatinine 145 mmol/l (N = 0.04 g/l). The 24 hour excretion of protein in the urine was 3.2 gm.

Immunoelectrophoresis of the urine revealed free kappa light chains.
Figure 1 — Contrast-enhanced CT scan. A) Showing bony erosion of sphenoid wing and temporal bone with extension into middle cranial fossa (M). B) Magnified view of orbits and sphenoid wings. Solid arrow points to destroyed lateral and medial orbital walls compared to normal left side (open arrow).

Bone marrow aspirate demonstrated diffuse infiltration by plasma cells. Cytoplasmic immunofluorescent staining confirmed IgD kappa type myeloma (Figure 4).

DISCUSSION

Neurological complications in multiple myeloma are relatively frequent, the most common being extradural spinal cord compression (Brenner et al., 1982; Silverstein and Doniger, 1963). According to Clarke (1954), cranial or intracranial myelomas can be broadly classified into three clinical groups:

(I) Syndromes of cranial nerve palsies — unilateral or bilateral VI, VII and unilateral I (Woodruff and Ireton, 1982).

(II) Intracranial tumour syndromes with or without involvement of skull bones (Weiner et al., 1966; Stark and Henson, 1981).

(III) Intra-orbital tumour syndromes (Clarke, 1953).

The present case appeared to be a myeloma arising from the body of the sphenoid wing, with extensive destruction of the nearby bone structures and compression of the right temporal lobe. The clinical picture was that of a retro-orbital mass lesion which, unlike most meningiomas, had produced extensive bony destruction. Bilateral orbital infiltration by myeloma has been reported by Hayes et al. (1980) in a patient with IgG myeloma and by Trecan et al. (1977) in a patient with IgM myeloma.

IgD myeloma (Fahey and Carbine, 1968) had been estimated to comprise about 2% of all myelomas (Jancelewicz et al., 1975). There is a male predominance (Fishkin et al., 1970) and 65% of 133 patients reported were younger than 60 years of age at diagnosis (Jancelewicz et al., 1975). As compared to IgA, IgG, and IgM myelomas, 90% of the IgD myelomas are of the lambda type (Pruzanski and Rother, 1970). They have a higher
incidence of extra-osseous spread, and are frequently associated with severe anemia, azotemia and amyloidosis (Jancelewicz et al., 1975). The mean survival is 13.7 months from diagnosis, and deaths are usually due to infections or renal failure.

Unilateral proptosis was observed in 6 of 133 patients with IgD myeloma, and in one of these patients there was associated ocular muscle paralysis (Jancelewicz et al., 1975). IgD myeloma has also been reported in a patient with sphenoid sinus involvement (Humphrey et al., 1983) and in two patients with meningeal infiltration associated with increased CSF IgD/albumin ratio (Schulman et al., 1980).

The present case is unusual in that the patient is female, older than age 60, and the paraprotein light chain is of the kappa type. As yet, there is no clinical evidence of amyloidosis, and she has only slight anemia and mildly impaired renal function. The difference in prognosis between patient with IgD myeloma exhibiting kappa vs. lambda light chains is as yet unknown. In general, patients with myeloma of the kappa chain type appear
to have a slightly longer mean survival time, mean 28.3 years vs. 18.3 years for the lambda type (Jancelewicz et al., 1975).

This case serves to illustrate that retro-orbital multiple myeloma should be considered in the differential diagnosis of patients presenting with painful unilateral proptosis.

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REFERENCES


