Acute fever and delayed leukoencephalopathy following low dose intraventricular methotrexate

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SUMMARY Nine out of 14 patients treated with intraventricular methotrexate (MTX) for meningeal carcinomatosis from breast carcinoma and surviving more than 4 months developed disseminated necrotising leukoencephalopathy (DNL). All four patients who had received both intraventricular MTX and whole brain radiotherapy developed DNL. Five of the six patients who experienced an acute febrile reaction with mild encephalopathic signs following intraventricular administration of MTX developed DNL after a mean time of 5 months and a low mean dose of 44 mg MTX. DNL was also noted in two patients without a previous febrile reaction or whole brain radiotherapy, following prolonged intraventricular MTX therapy after a mean time of 19-5 months and a mean dose of 147 mg MTX. These findings confirm the hazards of (1) high cumulative doses of intrathecal MTX and (2) combined intrathecal chemotherapy and whole brain radiotherapy. This study also suggests a possible relationship between an early and transient febrile reaction during intraventricular administration of MTX and the development of DNL.

Intrathecal (IT) methotrexate (MTX) is widely used to treat or prevent meningeal leukaemia and lymphoma.1 2 It is also successfully applied in patients with meningeal carcinomatosis, particularly from breast carcinoma.3-5 Treatment is usually combined with cranial or craniospinal radiotherapy and occasionally with systemic MTX.

The neurological complications of intrathecal MTX administration can be divided into acute and late effects. The acute effects develop within hours and consist of a transient aseptic meningitis or, rarely, a myelopathy or encephalomyelopathy.6-8 The reported frequency of MTX meningitis varies from uncommon to more than 50%.9-11 The symptoms of headache, vomiting and fever usually last 1 or 2 days. There is nuchal rigidity, usually associated with a modest CSF pleiocytosis and elevated CSF protein.9 The cause of this reaction is unknown and a number of factors have been implicated.6 11 12 Probably MTX meningitis is caused by the drug itself, as it is usually accompanied by elevated CSF MTX levels.13 Active meningeal infiltration by malignant cells may be a predisposing factor.13 14 The most important delayed toxic effect is disseminated necrotising leukoencephalopathy (DNL). It develops between 3 to 15 months after MTX administration and is clinically characterised by an insidious onset of personality changes, lethargy and dementia, usually followed by hemiplegia or quadripareisis and coma.9 15

Histological examination reveals areas of demyelination, axonal degeneration, astrocytosis and coagulative necrosis, predominantly in the periventricular white matter with or without vascular changes.16 17 This appears on CT as periventricular areas of decreased density. This complication almost exclusively occurs when IT MTX is combined with cranial irradiation or systemic MTX and seems to be dose-dependant.6 9 17 The incidence has been reported to be as much as 45% after high dose radiotherapy (>35 Gy) in combination with a larger dose of IT MTX.18

Hitherto only two patients have been described with DNL following IT MTX without additional cranial radiation or systemic MTX.19 20 The administered total dose of MTX in these patients was 217 mg/m² and 195 mg/m² respectively. The approximate risk of DNL due to IT MTX without whole brain radiotherapy or systemic MTX is assumed to be less than 1%, provided that a total dose of at least 50 mg of MTX has been administered.21

The present report analyses the occurrence of DNL in patients treated with IT MTX for meningeal car-
cinomatosis from breast carcinoma. It shows that the risk of DNL after single treatment with IT MTX may be considerable and that DNL may even occur after a total dose of less than 50 mg MTX. The role of possible contributing factors is investigated.

Patients and methods

Thirty-five patients with meningeal carcinomatosis from breast carcinoma were treated with the intraventricular administration of MTX via an Ommaya reservoir. In every patient the diagnosis was confirmed by the demonstration of malignant cells in the CSF. CT of the brain was performed in all patients before the insertion of the Ommaya reservoir. Intraventricular MTX therapy was only administered if concurrent systemic metastases were expected to be treatable. MTX was administered according to a protocol which has been described elsewhere: \(^5\) 5 mg of preservative-free isotonic MTX was injected whenever CSF MTX levels tended to fall below 10–9 mol/ml until the disappearance of tumour cells from the CSF. Thereafter, the interval of MTX administration was gradually increased to a maintenance programme of 5 mg once every 6 weeks. During the first treatment weeks in every patient CSF samples were taken at least twice a week from the Ommaya reservoir for culture, cytology, MTX levels and biochemical analysis.

Oral leukovorin was given after IT MTX if bone marrow suppression occurred. Fourteen of the 35 patients survived longer than 4 months after initiation of IT MTX therapy. These 14 patients were analysed for the occurrence of MTX neurotoxicity (table). The mean age of these patients when the diagnosis meningeal carcinomatosis was made was 53 years (range 39–72 years). In an attempt to detect factors which may contribute to the development of neurotoxic complications the area and the intensity of meningeal involvement were investigated, according to clinical signs, neuroradiological and CSF findings. Four patients received whole brain radiotherapy. Two patients (Nos 1 and 2) with parenchymal metastases received 3900 cGy and 4200 cGy respectively 4 and 17 months prior to IT MTX administration. The other two patients (Nos 3 and 4) received whole brain radiotherapy with respectively 1750 cGy and 3000 cGy, 6 weeks after the initiation of IT MTX. Patient 4 was irradiated because of a parenchymal metastasis. Patient 3 received additional craniospinal irradiation after IT MTX had been stopped because of possible MTX toxicity. During IT MTX therapy CSF had become negative for tumour cells, although simultaneously some clinical deterioration was noted. Radiotherapy had to be stopped in this patient at 1750 cGy (in 12 fractions) because of severe bone marrow suppression. The CSF remained free of tumour cells.

Two patients (Nos 6 and 11) received local radiotherapy to the skull base because of osseous metastases with 1150 cGy (in four fractions) and 3000 cGy (in 12 fractions) respectively 3 and 6 months before IT MTX treatment. Patient 11 received additional local radiotherapy to the hypophyseal area (2100 cGy in seven fractions).

The diagnosis DNL was made on the basis of typical clinical signs and CT findings. The clinical picture consisted of lethargy, ataxia, dementia and in some cases hemiparesis. CT showed the characteristic hypodensity of the periventricular white matter. Infectious meningitis or an exacerbation of the meningeal carcinomatosis were excluded. In three patients (Nos 4, 5, 7) the CSF Myelin Basic Protein (MBP) level was measured by a radio immunoassay technique (Dr. Out, Academic Medical Centre, Amsterdam) at the diagnosis of DNL. Postmortem examination was performed on two patients with DNL (patients 1 and 5).

Results

Acute fever (table)

A febrile reaction in the first weeks of IT MTX therapy occurred in six of the 14 patients. Temperature gradually rose 5 to 14 days (mean 9 days) after initiation of IT MTX treatment, that is, after the second or third intrathecal injection and lasted 6 to 16 days (mean 11 days). In most cases fever was mild with temperatures between 38.5°–39°C and only occasionally rising to 40°C. Accompanying symptoms and signs were also mild and consisted of headache, nausea and dizziness and in some cases a slight ataxia.

Table Summary of treatment and its neurotoxic complications in patients with meningeal carcinomatosis from breast carcinoma

<table>
<thead>
<tr>
<th>Patient/Age (yr)</th>
<th>Dose of IT MTX</th>
<th>Whole brain irradiated</th>
<th>Acute fever</th>
<th>DNL</th>
<th>Onset of DNL</th>
<th>Survival time after treatment</th>
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<tr>
<td>1/48</td>
<td>55 mg</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>5 m</td>
<td>6 m</td>
</tr>
<tr>
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<td>–</td>
<td>+</td>
<td>7 m</td>
<td>7 m</td>
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<td>+</td>
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<td>–</td>
<td>8 m</td>
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<td>+</td>
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</tr>
</tbody>
</table>

DNL = Disseminated necrotising leukoencephalopathy.
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Nuchal rigidity did not increase in any of these patients. All cultures were negative. In all patients fever disappeared spontaneously without antibiotic treatment. In four patients (Nos 4, 5, 7, 10) IT MTX was continued throughout the period of fever. Only in one patient (No. 3) was some relation apparent between the administration of MTX and the fever shown by temperature peaks within 24 hours following IT MTX.

No significant difference was seen in the extent, the intensity or the course of the meningeal involvement between patients with and without fever. During the febrile reaction the ventricular CSF cell count and protein concentration remained at the same level or became within normal limits.

CSF MTX concentrations in the toxic range as defined by Bleyer et al13 were measured at one or more occasions during the first 4 weeks of IT MTX treatment in four of the six patients with a febrile reaction, and in three of the eight patients without a febrile reaction (fig 1). Notably, no difference was noted in the CSF MTX levels during and after the period of fever in the four patients in whom IT MTX administration was continued. In three patients IT MTX was changed into cytosine arabinoside (ARA-C), given in a dose of 40 mg in 2 ml of preservative-free water. In Patient 4 a total dose of 160 mg ARA-C was given because of simultaneous whole brain radiotherapy for parenchymal metastases. Patient 6 received 40 mg ARA-C because of the febrile reaction during IT MTX. As fever persisted ARA-C was also discontinued. Patient 8 received 80 mg ARA-C because of a clinical and cytological response failure to IT MTX.

Leukoencephalopathy (table)

Nine of the 14 patients developed DNL 3 to 26 months after the initiation of IT MTX. All four patients who had received whole brain radiotherapy (Nos 1, 2, 3, 4) developed DNL 4 to 8 months (mean 6 months) after the start of IT MTX. The mean total dose of IT MTX in these patients was only 36 mg (range 25–55 mg). Both patients who received whole brain radiotherapy after IT MTX experienced a febrile reaction in the first weeks of IT treatment. One of these patients (No. 3) received only 1750 cGy (in 12 fractions) after a total dose of 35 mg MTX. MTX was stopped before whole brain radiotherapy. In the other patient (No. 4) MTX was changed into ARA-C during radiotherapy for parenchymal metastases because of the risk of provoking DNL. She eventually received 30 mg MTX and 160 mg ARA-C IT. Two months after radiotherapy she developed signs of DNL. CT showed disappearance of the parenchymal metastasis as well as of the contrast enhancement along the cerebral sulci but on the contrary periventricular hypodensity. Five of the 10 patients without whole brain radiotherapy developed DNL after treatment with IT MTX. The mean total dose of IT MTX in patients with DNL was 90 mg (range 15–164 mg) versus 61 mg (range 30–100 mg) in those without DNL. Three of the five patients with DNL had experienced a febrile reaction in the early stage of IT treatment. A mean dose of only 52 mg IT MTX (range 15–85 mg) had been given in these three patients and DNL developed after a short mean time of 5 months (range 3–7 months) (fig 2). In contrast, in the two patients who developed DNL without a previous febrile reaction DNL was diagnosed at 13 and 16 months, respectively after 130 and 164 mg IT MTX. No relation was found between the CSF MTX levels during the first 4 weeks of intensive IT MTX administration and the development of DNL (fig 1). In addition, no relation could be found between the development of DNL and the area or intensity of malignant infiltration. The CSF MBP level (normal <1 ng/ml) was 2.8 ng/ml in Patient 4, 15 ng/ml in Patient 5 and 5.5 ng/ml in Patient 7.
The mean survival time after the initial diagnosis of DNL was 5 months (range 0–16 months). One patient (patient 7) is still alive 4 months after the diagnosis of DNL.

All patients showed a steady deterioration of the DNL. In all patients DNL was the major cause of death or an important contributing factor. However, the mean survival time from the start of IT MTX therapy was 14 months in the patients who developed DNL in contrast with 7 months in those without DNL.

Neuropathological examination showed macroscopically visible foci of necrosis and demyelination in the cerebral periventricular white matter in both patients with DNL examined (fig 3). In Patient 5 these foci were predominantly located in the direct vicinity of the ventricles.

For microscopic examination haematoxylin and eosin, Kluver, PAS and Kossa stains were used. In both patients tumour cells were found in most parts of the subarachnoid space. In Patient 1 a section of the right frontal area showed necrotic lesions, surrounding small areas with relatively intact white matter. In the necrotic lesions macrophages were seen and fields of astrocytosis. The vessels in these areas were thickened by fibrinoid necrosis, with hyalinisation and narrowing of the lumen without real occlusion. The Kossa stain was negative. Microscopic examination of both frontal areas in Patient 5 demonstrated necrotic areas in the white matter with myelin degeneration, cellular remnants, macrophages and some glial reaction and vascular proliferation (fig 4). Hyalinoid and fibrinoid-necrotic changes of blood vessel walls were seen with narrowing of the lumen occasionally with complete occlusion. Amyloid was not observed. The Kossa stain for calcium in the vessel walls was slightly positive.

Discussion

The acute febrile reaction following intrathecal administration of preservative-free MTX in six patients in the present study differs clearly from MTX meningitis. Fever gradually developed after the second or third dose of MTX without an obvious relation to the moment of injection while it lasted much longer than 1 or 2 days. There were in fact no signs of a meningeal reaction. Nuchal rigidity was absent and CSF did not show a cellular reaction or elevated protein. No significant relation was found with elevated CSF MTX levels, nor with the area or intensity of the malignant meningeal involvement.
The dose of MTX per injection in our patients was lower than usually applied.\textsuperscript{3,4,6,11} DNL almost exclusively occurs when IT MTX is combined with whole brain radiotherapy and/or systemic MTX.\textsuperscript{9,17} The incidence of DNL increases with the cumulative dose of IT MTX\textsuperscript{17} and may be correlated with elevated CSF MTX levels.\textsuperscript{22} The present study shows a clear relation between DNL and (1) whole brain radiotherapy, (2) the cumulative dose of MTX and (3) an early febrile reaction. No close relation was found between the occurrence of DNL and ventricular CSF MTX levels. The patients with a febrile reaction developed DNL after a short interval and following a low dose of MTX. Two of these patients had also received whole brain radiotherapy. However, as in both patients IT MTX had been stopped before radiotherapy it is questionable whether radiotherapy has substantially contributed to the development of DNL.\textsuperscript{9,14} Moreover, the total dose of 1750 cGy in 12 fractions in one of these patients is lower than the estimated minimal neurotoxic dose.\textsuperscript{6,17}

Only two cases of DNL following IT MTX without radiotherapy or systemic MTX have been previously described.\textsuperscript{19,20} The cumulative dose of IT MTX of 217 mg/m\textsuperscript{2}\textsuperscript{19} and 195 mg\textsuperscript{20} is in the range of the total dose of MTX (130 mg and 164 mg) that was given in the two patients in the present study who developed DNL without a previous acute febrile reaction. In other reports concerning DNL following IT MTX the patients had also received high dose intravenous MTX or partial or whole brain irradiation.\textsuperscript{16,22-24} This apparent rarity of DNL after IT MTX alone may be partly explained by the fact that in most treatment regimens for neoplastic meningitis IT MTX is combined with other treatment modalities.

Elevation of MBP in CSF is strongly correlated with active demyelination and is likewise always encountered in DNL.\textsuperscript{25,26} Accordingly, in the present study CSF MBP was elevated in patients with DNL, including patients treated with IT MTX without whole brain radiotherapy.

The pathogenesis of these various types of MTX toxicity is obscure. In meningeal carcinomatosis CSF flow abnormalities may occur in 70% of the patients. These abnormalities are often transient during the first weeks of treatment and may consist of ventricular obstruction as well as of a flow delay over the cortical convexities. Importantly, these CSF flow disturbances are not closely associated with abnormalities in CSF cell count or protein nor with CT findings.\textsuperscript{27} So the early transient encephalopathic symptoms in our patients might have been caused by an increased MTX diffusion into the cerebral parenchyma due to
an early transient decreased CSF flow.

For the development of DNL following MTX several mechanisms have been considered. Apart from the direct toxic effect of MTX on glial and endothelial cells, the mental changes have been related to deficiencies of folic acid, serotonin and biogenic amines, caused by MTX. The neurotoxic effect of MTX is potentiated by cranial irradiation which induces endothelial damage resulting in an increased diffusion of MTX into the brain parenchyma.

The predilection for the periventricular white matter has been explained by impaired CSF flow from the ventricles with increased transependymal diffusion. Impaired flow over the convexities with increased absorption from the deep Virchow-Robin spaces may be a contributing factor. The absence of DNL after prophylactic treatment with IT MTX is in agreement with the significance of CSF flow disturbances due to the neoplastic meningitis. Postmortem examination in Patient 5 showed predominant demyelination and necrosis in the direct vicinity of the lateral ventricles suggesting that a pathologic transependymal diffusion of MTX may also occur at so-called non-toxic ventricular CSF MTX levels.

A sensitisation or hypersensitivity response, as proposed in some other cases of MTX neurotoxicity, can not be excluded as an underlying mechanism in causing the acute febrile reaction and DNL after low dose of MTX. However, disappearance of the fever during continued MTX administration is not in accordance with this hypothesis. ARA-C may cause DNL after high dose IV infusion but not after intrathecal administration and therefore did not contribute to the development of DNL in our patients.

The neuropathological findings show that apparent irradiation specific vascular changes can also be produced by IT MTX alone. Histologically, there was no striking difference between the two examined patients, although in Patient 1 the lesions can be interpreted as more acute. This patient had experienced a rapidly progressive course of the disease and only lived 1 month after the first signs of DNL. It may be speculated whether previous whole brain radiotherapy in this patient has played a role in the rapid development of these extensive necrotic lesions.

This study shows that DNL is not a rare complication of IT MTX therapy. The incidence is related to the cumulative dose of MTX and the duration of treatment as well as to whole brain radiotherapy. In addition, DNL may be induced by less than 50 mg IT MTX in patients who have developed an acute reaction to IT MTX that consists of fever and mild
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encephalopathic symptoms. Because of this serious and debilitating late complication one has to be aware of this possible relation. As DNL occasionally is reversible it might be advisable to withhold MTX as soon as this acute encephalopathic reaction is noted, and replace it by ARA-C. The surprisingly high incidence of this serious complication makes the development of other chemotherapeutic agents or treatment modalities for meningeval carcinomatosis more urgent.

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References