Ischaemic neuropathy of the lumbosacral plexus following intragluteal injection

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SUMMARY A lesion of the lumbo sacral plexus may result from an inadvertent intra-arterial injection of vasotoxic drugs into one of the gluteal arteries. Symptoms and follow-up of three cases are reported. The neuropathy is attributed to a toxic endarteritis with retrograde propagation of spasm and thrombosis. Swelling and bluish discoloration of the buttocks ("embolia cutis medicamentosa") as well as an impaired circulation in the homolateral leg are associated with the neurological syndrome in fully developed cases and makes possible a correct diagnosis.

The obstruction of a main artery by embolism, thrombosis or following trauma may lead to ischaemic neuropathy. The neurological syndrome, predominantly sensory impairment and (more rarely) paralysis, or in milder cases pain and paraesthesia, depends upon the duration of ischaemia and collateral blood supply.1-3 If the ischaemia is of short duration a rapid and complete recovery of nerve functions will occur in most cases, but ischaemia of a longer duration can lead to permanent structural damage.2 4 Since the oxygen turn-over of the peripheral nerve is rather low and blood supply is abundant, ischaemic neuropathies will rarely occur if only one of the main arteries or only some, but not all of the distal branches of the nerve vessels are obstructed. However, partial or total segmental necrosis of nerves regularly occurs if microembolism, microthrombosis or spasms obstruct large portions of the epineural and perineural plexus.5 8 This mechanism is responsible for cases of neuropathies after intra-arterial injection of angiototoxic substances, for example into the brachial artery,6 7 the inferior gluteal artery9-10 and the umbilical artery of the newborn.11 12

An inadvertent injection of vasotoxic drugs into one of the gluteal arteries may also result in a toxic angioopathy with ischaemic damage of the sciatic or gluteal nerves. With the retrograde propagation of angiospasms and thrombosis, the homolateral lumbosacral plexus can suffer ischaemic damage, as observed in three cases.

CASE REPORTS

Case 1 (BF, a 42 year old male): an injection of about 4.5 ml Ultrademoplus (500 mg Phenylbutazon-Natrium, 500 mg Aminophenazon, 45 mg Lidoceain-HCl, 4-5 mg Dexamethason, 9 mg Prednisolon and 2-8 mg Cyanocobalamin) was given into the left buttock because of lower-back pain. Immediately a severe local pain was felt, lasting for about 30 s. Ten to fifteen minutes later, the patient noticed numbness in the left foot which crept gradually upwards. Half an hour after the injection, standing and walking were no longer possible because of a severe pain as well as a weakness in the left lower limb. The neurological examination revealed a nearly complete paralysis of the left leg, including the hip-flexors and the gluteal muscles; deep tendon reflexes were absent. There was severe sensory impairment in the territories of the femoral, lateral femoral cutaneous and common peroneal nerves, with less marked sensory disturbances in the other areas of the left lower limb. There was painful swelling and bluish discoloration of the left buttock. Additional swelling at the left paravertebral region above the pelvis ("Embolia cutis medicamentosa," fig 1) occurred and was followed by a deep gangrene within a few days. The left foot appeared colder and livid. A few days later, the persistent pain in the left leg was intensified by paroxysms. On
the fifth day, the intermittently slightly improved weakness deteriorated again with one severe paroxysm of pain. In addition, micturition was intermittently disturbed and painful. The patient was not incontinent and defaecation was normal. The electromyographic examination revealed complete denervation of the quadriceps muscle and partial denervation of the other muscle-groups of the lower left limb, as well as of the paravertebral muscles at the L4/L5 level. The ninhydrine-test indicated a hypohidrosis of the left foot.

General examination and blood-tests revealed no signs of diabetes mellitus or general vascular disease. Myelography was normal. The CSF protein was increased to 70 mg/dl; further CSF examinations were normal.

One year later, pain had considerably decreased, but was still present, mainly in the anterior aspect of the thigh, and was aggravated by walking. Neurological examination revealed severe paresis of the extensors of the knee, foot and toes, moderate paresis of the hip flexors and slight paresis of the other muscle-groups (including the gluteal muscles). There was moderate hypeaesthesia combined with hyperpathia within the territories of the femoral, lateral femoral cutaneous and common peroneal nerves. Arterial pulses of the femoral and popliteal arteries were slightly weaker on the left side. The skin temperature of the left foot was reduced by 1°C. The oscillogram showed signs of a compensated arterial obstruction. Diagnosis: Ischaemic neuropathy of the left lumbosacral plexus with electromyographical signs of root involvement.

Case 2 (WH, a 38 year old male): One ampoule Delphimix (600 mg Phenylbutazon-Natrium, 6 mg Cinchocain, 40 mg Triamcinolon-diacetat, 1 mg Cyanocobalam) was injected into the left buttoc, because of lower back pain. Five to six hours later the patient awoke from sleep because of a numbness in the left leg. Another six hours later he experienced severe waxing and waning pain predominantly at the anterior aspect of the thigh and he was no longer able to stand. The patient was admitted to the hospital, where a painful swelling and bluish discoloration of the skin were observed at the upper part of the buttoc. There was a severe paresis of the flexors and adductors of the hip and of the extensors of the knee, as well as marked hypeaesthesia and hyperpathia anteromedially at the thigh were found. For several weeks pain was almost totally resistant to analgesic drugs. Pain disappeared only slowly in the course of the next two years. General examination and blood tests were normal.

A follow-up examination two years later
revealed slight residual paresis of the above mentioned muscle groups as well as marked hypaesthesia on the medial aspect of the knee. Knee-jerk was diminished and adductor reflex was absent. The electromyographical investigation showed signs of partial denervation of the iliopsoas, adductor longus and rectus femoris muscles. No abnormalities were found in the paravertebral muscles (L3/L4). Diagnosis: Ischaemic neuropathy of the left lumbar plexus.

Case 3 (BA, a 13 month old male): An injection of penicillin into the left buttock was followed by a painful local swelling, and a flaccid paralysis of the left lower limb. The neurological examination 4-5 months later revealed a still complete paresis of the whole leg, except for a weak residual innervation of the toe-flexors (fig 2). Deep tendon reflexes were absent in the left, but were present in the right leg. There was no reaction to painful stimuli in the left foot and lower leg.

EMG examination and electrical nerve stimulation showed nearly complete denervation of the long and short flexors of the toes, as well as complete denervation of the gluteal, quadriceps, tibialis anterior and soleus muscles. There were no signs of denervation in the paraspinal muscles (L3-L5). General examination, blood tests, CSF and X-rays of the pelvis and the spine were found to be normal. Diagnosis: Ischaemic neuropathy of the left lumbosacral plexus.

Discussion

Lesions of the lumbosacral plexus following an intragluteal drug injection cannot be explained by a direct mechanism, such as toxic neuropathy of the sciatic and gluteal nerves. The damage, because of its distance from the site of injection, suggests the importance of a vascular factor. An inadvertent injection of vasotoxic or crystalline drugs into the inferior gluteal artery, which supplies the proximal segment of the sciatic nerve, is known occasionally to result in a sciatic nerve lesion.8 9 13 The underlying cause is most likely to be either embolic obstruction by crystals14 or a toxic endarteritis with spasms and thrombosis, which spread to the epineural and perineural blood vessels and cause segmental infarction.

Since the skin of the buttock is largely supplied by the same artery, it is not surprising that these particular lesions of the sciatic nerve are invariably associated with a painful swelling and a bluish discoloration of the buttock, which is then sometimes followed by gangrene ("embolia cutis medicamentosa").8 10 13 14

A similar mechanism could be responsible for lumbosacral plexus lesions, if a retrograde propagation of spasm and thrombosis occurs. Some observations are in favour of such a possibility. Homolateral gangrene of rectum, bladder, penis, scrotum and vagina have been described following an intragluteal injection8 13 15 and suggest a circulatory disturbance within the distribution of the internal iliac artery as their possible cause. Provided that a still more extensive retrograde propagation of the toxic angioptathy occurs, the external iliac artery may be occluded in addition, with subsequent ischaemia of the lower limb.16 17

Since the lumbosacral plexus receives its blood supply from branches of the iliac arteries (fig 3) it is reasonable to assume that lumbosacral plexus lesions may be another consequence of such a wide-spread toxic angioptathy. In one of our cases (BF) a deep gangrene of the skin above the pelvis (fig 1) developed within the distribution of the fourth lumbar artery. This artery anastomises with the iliolumbal artery. The circulatory disturbance even in this proximal branch of the internal iliac artery suggests a widespread retrograde propagation of the toxic angioptathy.
obturator nerve palsy. Paresis invariably followed an injection into the ipsilateral buttock and was accompanied by immediate severe pain, discoloration of the buttock and a circulatory disturbance in the leg.

Regarding the distribution of the neurological signs in these and our own cases we can differentiate three patterns of plexus damage, involving (1) the lumbar plexus or its femoral branch, or (2) the lumbar and sacral plexus, or (3) the lumbosacral plexus including some lumbosacral nerve roots (Table). In very rare instances, the toxic angiopathy following an intraarterial injection in gluteal arteries can spread to the aortic bifurcation and result in a flaccid paraplegia.\textsuperscript{19}\textsuperscript{20}

Table Various neurological syndromes following an inadvertent drug injection into one of the gluteal arteries

<table>
<thead>
<tr>
<th>Distribution of the toxic angiopathy</th>
<th>Involved nervous structure</th>
<th>Neurological syndrome</th>
<th>Author</th>
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<tbody>
<tr>
<td>Inferior gluteal artery</td>
<td>Sciatic nerve</td>
<td>Sciatic or peroneal nerve palsy</td>
<td>Gammel (1928)</td>
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<tr>
<td>Ipsilateral iliac arteries</td>
<td>(a) Lumbar plexus</td>
<td>Femoral- or femoral and obturator nerve palsy</td>
<td>Vasilescu and Stamatoiu (1965)</td>
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<td></td>
<td>(b) Lumbar and sacral plexus</td>
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<td>(c) Lumbosacral Flaccid plexus with root involvement</td>
<td>Flaccid paraplegia of the ipsilateral leg</td>
<td>Stöhr et al (Case 3)</td>
</tr>
<tr>
<td>Aortic bifurcation</td>
<td>Spinal cord</td>
<td>Flaccid paraplegia</td>
<td>Heckenroth (1955)</td>
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<td></td>
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<td>Merling (1959)</td>
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The ischaemic plexopathies following intra-arterial injection in gluteal arteries exhibit striking similarities with a syndrome in newborns, which results from an injection of vasotoxic drugs into the umbilical artery. The umbilical artery is a branch of the internal iliac artery, as are the gluteal arteries (fig 3). In this case the injection may result in a discoloration and sometimes gangrene of the buttock and in a sciatic nerve palsy. Occasionally a femoral nerve palsy and a gangrene of intrapelvic structures are associated.\textsuperscript{11}\textsuperscript{12}\textsuperscript{21}\textsuperscript{22} As in lumbosacral plexopathies following an intragluteal injection, the development of the neurological signs may be delayed in newborns.\textsuperscript{23}\textsuperscript{24} There is, however, one striking difference in the pattern of nerve damage following intragluteal and intrarumbral injections. When there is a drug injection into the umbilical artery of the newborn, there is a predominant involvement of the sciatic nerve. This is probably
due to the greater functional importance of the inferior gluteal artery at this age. Plexus lesions following an injection into the buttock however case lesions of the lumbosacral plexus, especially concerning its femoral nerve portion. The same pattern has frequently been observed in postoperative plexus-lesions, where the stretching of nerves and nerve vessels appears to be the most damaging factor. The possibility of successful therapy of vascular occlusion due to intra-arterial injection is greater when quickly diagnosed. Intra-arterial injection may be suspected if the patient immediately complains about severe local pain and may be ascertained by aspiration of arterial blood before retracting the needle. In this situation, 5 to 10 ml of Procaine 1% (without Suprarenin) and antispasmodic drugs such as Papaverine (40–80 mg in 10–20 ml isotonic saline solution) may prevent the arterial spasms. In order to secure the retrograde propagation of these drugs the pressure of injection should be high. Most frequently the correct diagnosis will not be reached before the retraction of the needle. In these cases Papaverin should be given intravenously. In addition, antithrombotic therapy with heparin and sympathetic blockade are recommended.

Intra-arterial injection into one of the gluteal arteries is more likely and if it happens more hazardous if the injection is performed on the medial aspect of the buttock, near the exit of the gluteal arteries from the pelvis. Therefore an injection into the upper and outer part of the buttock is recommended not only to prevent a direct sciatic nerve lesion, but also to diminish the possibility of an intra-arterial injection. Moreover, an aspiration test is mandatory prior to every injection into the buttock. Lumbosacral plexus lesions following an injection into the buttock within a few hours are a rather typical syndrome which can hardly be confused with any other syndrome. A coincidence of intragluteal injection and lumbosacral plexus lesion without causal relation seems already very unlikely, especially unlikely when the rarity of the non-traumatic plexus lesions is considered. Moreover, there should be some evidence for another cause, which occasionally leads to an acuteplexopathy, like diabetes mellitus, viral infections, haemophilia or anticoagulation therapy. Other signs of a widespread toxic angiopathy such as painful swelling and discoloration of the buttock, ipsilateral gangrene of intrapelvic structures and ischaemia of the lower limb support an ischaemic plexus-lesion following an intra-arterial drug injection.

References


24 San Agustin, Nitowsky HM, Borden JN. Neonatal sciatic palsy after umbilical vessel injection.

