

Involvement of Regional Lymph Nodes after Penetration of *Schistosoma mansoni* Cercariae in Naive and Infected Mice

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The parotid lymph nodes of naive and previously infected Balb/c mice were studied after, respectively, infection and re-infection with cercariae of Schistosoma mansoni via the ears. Schistosomula were able to pass through the lymph node by following the lymph flow or by penetrating the veins of the medullary cords. The number of nodal mast cells was higher from day 2 to 6 of primary infection; and from day 5 to 11 of re-infection. The amount of degranulating mast cells was significantly higher at day 4 of infection and at day 1 of re-infection. Eosinophils characterized the nodal inflammatory processes observed after day 5 in both primarily-infected and re-infected mice. However, only in the latter the eosinophils were able to adhere to the larval surface. In primarily-infected mice, no intranodal larva presented signs of degeneration. In contrast, in re-infected animals, some degenerating larvae were found inside eosinophilic infiltrates. The eosinophils reached the nodal tissue by migrating through the high endothelial venules and their collecting veins.

Key words: *Schistosoma mansoni* - lymph nodes - schistosomulum migration - eosinophils - mast cells - high endothelial venules

When cercariae of *Schistosoma mansoni* penetrate the host skin, they transform into schistosomula by undergoing several changes such as tail loss, discharge of secretory glands, glycocalyx shedding, and other surface alterations (Gordon & Griffiths 1951, Stirewalt 1974). On their way to the dermal and subdermal layers, the schistosomula may spend several days till their passage to blood and/or lymph vessels at the dermal-hypodermal junction. By incubating slices of lymph nodes, Miller and Wilson (1978) demonstrated that schistosomula can be recovered from mouse nodes up to day 8 of infection, with peak at day 4. Histological studies have shown that schistosomula leave the regional lymph nodes of primarily-infected mouse, passing through the lymphatic sinuses to reach the efferent lymphatic (Wheater & Wilson 1979). However, the fate of schistosomula that reach lymph nodes after re-infection remains to be investigated.

Cercarial penetration induces an inflammatory

process (Colley et al. 1972), the skin thickness attaining the maximal values 24-48 hr after infection (Lichtenberg et al. 1976, Incani & McLaren 1984). However, the host leukocytic response is regarded as absent in regional lymph nodes of primarily-infected mice (Wheater & Wilson 1979). In infected mice, the changes subsequent to a challenge infection happen earlier, the skin thickness attaining maximal values in a few hours (Colley et al. 1972, Lichtenberg et al. 1976, Savage & Colley 1980, Bentley et al. 1981) but there is no study on regional lymph nodes. The behaviour of the nodal mast cells at both infection and re-infection is unknown.

The present work comprises a histological study of a regional lymph node after infection and re-infection of Balb/c mice with *S. mansoni* cercariae. It is centered on the nodal mast cells, inflammatory processes, and the fate of the schistosomula that reach the node. Previous studies have shown that the mouse ears are adequate for cercarial penetration and for studying the phenomena subsequent to the penetration (Gerken et al. 1986). Therefore, this was the site of stimulation and, consequently, the parotid lymph node that drains the ear was the object of study. Balb/c mice developed resistance to re-infection (reviewed by Dean 1983) even after immunization with irradiated cercariae (Sher et al. 1982). The resistance to challenge infection induced by the short-living attenuated larvae is both thymus and B lymphocyte dependent.

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MATERIALS AND METHODS

Cercariae - Cercariae of *S. mansoni* (LE strain, Belo Horizonte, Brazil) obtained from laboratory-reared *Biomphalaria glabrata* were concentrated as described by Pellegrino and Macedo (1955).

Mice and infection - Sixty-eight male Balb/c mice aged 60 days at the beginning of the experiments were used as: (a) 6 mice with 12-13 weeks of infection (infected mice); (b) 41 age-matched infected mice to be killed daily from day 1 to 8 and at day 11 of re-infection, (challenged mice); (c) 18 age-matched naive mice to be killed at days 2, 4, 6, and 8 after infection (primarily-infected mice); (d) 4 age-matched non-infected mice. Four to six animals were killed under ether anesthesia at each day of infection or re-infection, in two experiments each one with about half the animals.

For obtaining the infected animals, each mouse received a subcutaneous injection of 15 to 20 cercariae in 0.5 ml. For studying the response of lymph nodes to the primary infection and re-infection, anesthetized (Nembutal, 35 mg/kg) mice were infected percutaneously by the right ears with cercariae of *S. mansoni*. The animals were held on a wooden board containing holes in such a way that the right ears remained immersed in a suspension with 250-300 cercariae, for 1 hr (Gerken et al. 1986). On the average, 80% of the cercariae penetrated each ear.

Histological methods - The right parotid lymph nodes were fixed in 1% cetyl pyridinium chloride (w/v) in 10% (v/v) formaldehyde, for 48 hr at room temperature. After alcohol dehydration, the tissues were embedded in glycol methacrylate (JB-4, Polysciences), and serially sectioned at 5 μ m (Reichert Jung microtome, model 1140). The sections were alternately stained with Giemsa stain (Merck, São Paulo) and 0.5% toluidine blue (Sigma) containing 1% sodium borate.

Mast cells classed as intact and degranulating cells were counted at magnification X 400, in all Giemsa-stained sections at a 25 μ m interval. In each section, all mast cells exhibiting well-delineated nuclei were counted. An ocular disc containing a 5 mm square prevented recounting. The results were expressed as the mean number of degranulating or intact mast cells per lymph node section.

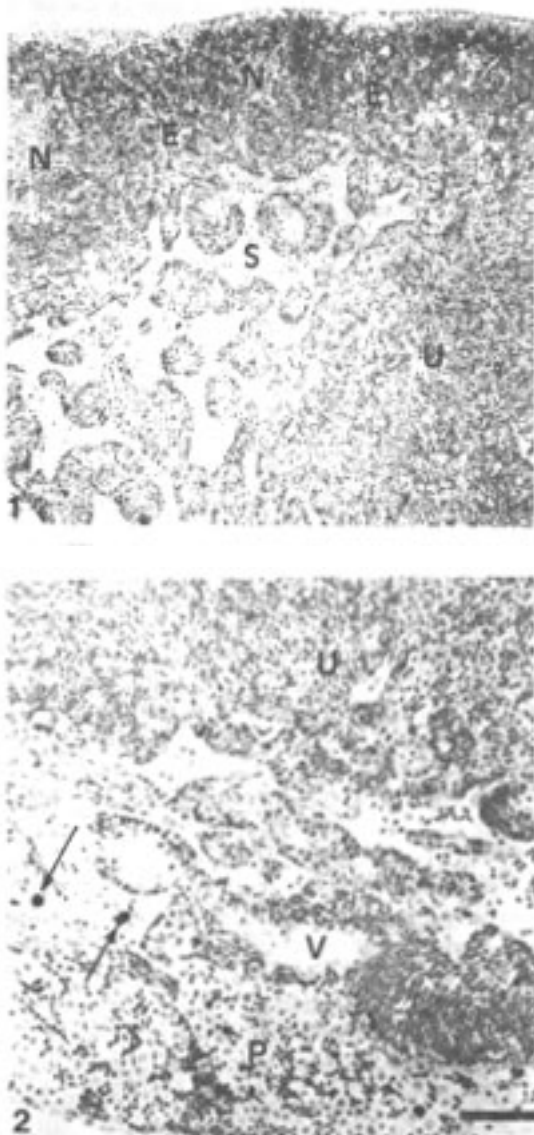
For estimating the number of schistosomula in each lymph node, all toluidine blue stained sections at a 25 μ m interval were used.

RESULTS

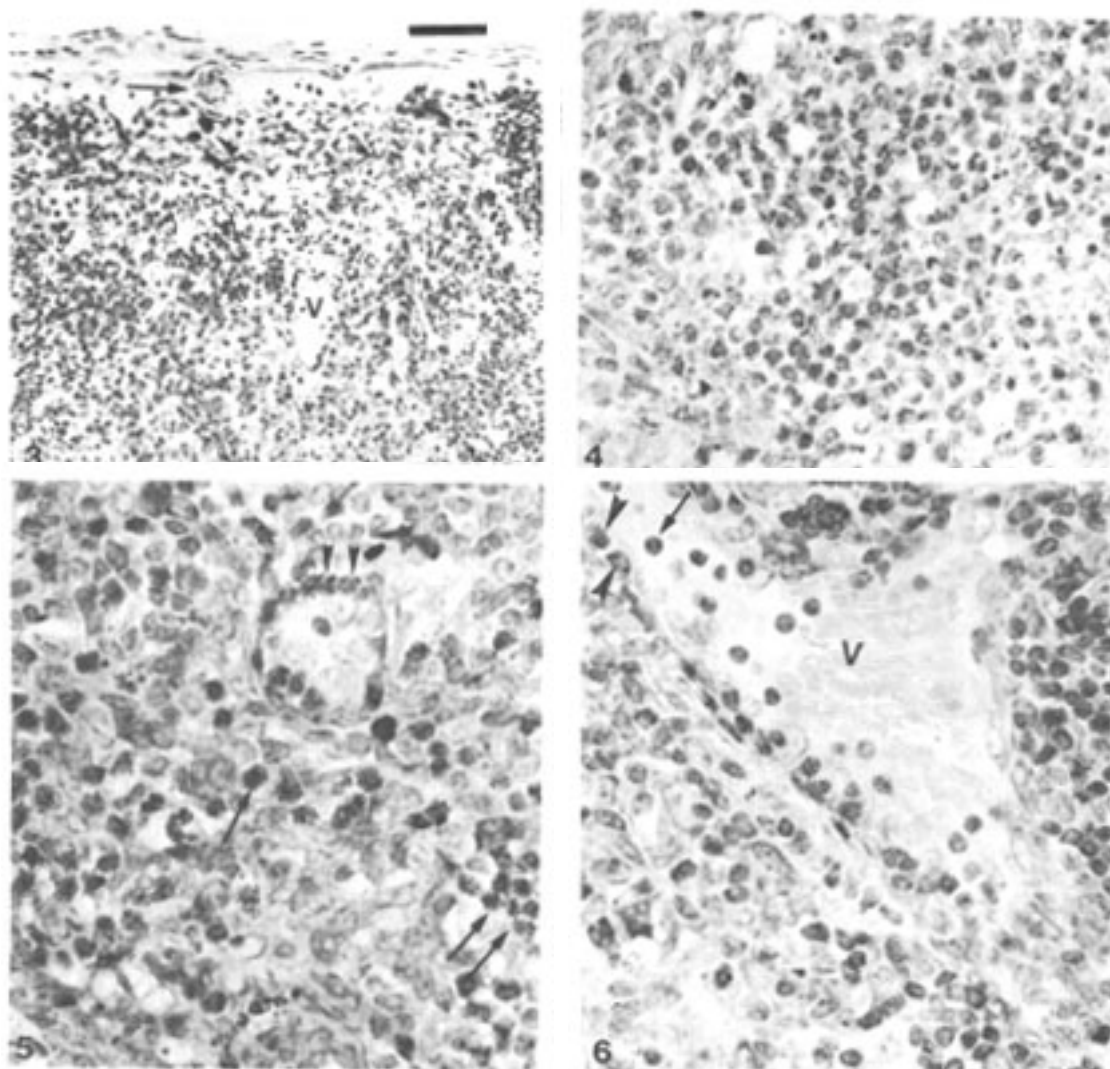
The parotid lymph node is a non-segmented node (Fig. 1), displaying a continuous peripheral cortex and two to three deep cortex units. An impressive finding was the abundance of macrophage like cells in the perihilar segments of the med-

ullary sinuses (Fig. 2). Mast cells occurred in small number, being found mainly at the perihilar segments of the medullary sinuses (Figs 2, 7). A few of them lay in the subcapsular sinus or just below it, usually in cortical extranodular zones (Fig. 3).

Mast cell number - The *S. mansoni* infection had no effect on the distribution of nodal mast cells either in naive or in infected animals. However, as



Histological appearance of mouse parotid lymph node (uninfected). Toluidine blue-borate. Fig. 1: the peripheral cortex showing lymphoid nodules (N), extranodular zones (E) and medullary tissue touching the cortical components, between deep cortex units (U). Fig 2: medullary sinuses and cords below a deep cortex unit (U) and in the perihilar region (P) in which the cords are thinner and the sinuses are plenty of macrophage-like cells. A medullary cord presents a collecting vein (V). Two mast cells are arrowed. Bar = 135 μ m (1); 48 μ m (2).



Parotid lymph nodes at day 6 of primary infection with cercariae of *Schistosoma mansoni*. Fig. 3: a transversal section of a schistosomulum located in the subcapsular sinus (arrows) and two intact mast cells just below (arrowheads). Granulocyte nuclei revealed an inflammatory process close to a high endothelial venule (v) in an extranodular zone. Toluidine blue-borate. Fig. 4: Giemsa stain shows eosinophils (the cytoplasm darker than the nucleus) in the inflammatory process of an extranodular zone. Fig. 5: a high endothelial venule displaying eosinophils in its lumen, wall (arrowheads) and neighboring extranodular tissue (arrows). Giemsa stain. Fig. 6: a medullary cord exhibiting several eosinophils; its vein (V) also shows eosinophils in the lumen (arrow) and wall (arrowheads). Giemsa stain. Bar = 48 μ m (3.6); 19 μ m (4.5).

summarized in Tables I and II, it induced marked changes in the amount of intact and degranulating mast cells. In primarily-infected animals, the number of intact mast cells (Table I) was higher from day 2 to 6 of infection, with peak at day 4. In challenged mice, a significant rise appeared only 5 days after cercarial stimulation, the values increasing till day 7, and remaining high at least until day 11. The number of mast cells presenting signs of degranulation (Table II) was higher at day 4 of infection and day 1 of re-infection. At day 6 of infec-

tion and from day 3 to 8 of re-infection, the number of degranulating mast cells was significantly lower as compared with the values found for uninfected and for infected but non-challenged animals. Degranulating mast cells had no relation to the presence of larvae, intact mast cells being frequently seen close to the larvae (Fig. 3).

Larval migration and inflammatory processes - Table III presents an estimate of the number of schistosomula in lymph nodes at different periods of the infection and re-infection. In primarily-in-

TABLE I

Mean number of intact mast cells per right parotid lymph node section in naive and infected Balb/c mice. Infection or re-infection with *Schistosoma mansoni* cercariae through the right ear. In parenthesis, the number of animals

Days after (re)infection	Naive mice	Re-infected mice
0	8.53±3.22 (4)	12.06±2.90 (6)
1	not done	9.70±2.05 (6)
2	15.99±0.55 ^c (4)	15.15±2.46 (4)
3	not done	13.79±3.30 (5)
4	29.22±11.76 ^b (5)	15.57±8.49 (4)
5	not done	27.97±11.33 ^a (4)
6	16.66±5.29 ^a (5)	32.61±10.96 ^c (5)
7	not done	34.57±8.73 ^c (4)
8	12.47±5.36 (4)	23.35±8.30 ^a (5)
11	not done	19.24±12.24 ^a (4)

Values are significantly different from the first value of the same column, when marked with *a*: (P<0.05), *b*: (P<0.005), or *c*: (P<0.001).

TABLE II

Mean number of degranulating mast cells per section of right parotid lymph nodes in naive and infected Balb/c mice. Infection or re-infection with cercariae of *Schistosoma mansoni*, through the right ear. Number of animals as in Table I

Days after (re)infection	Naive mice	Re-infected mice
0	4.30±1.58	7.74±1.19
1		12.67±4.09 ^a
2	6.09±4.35	8.53±1.35
3		5.90±1.37 ^a
4	10.57±2.27	4.30±2.30 ^a
5		4.99±1.00 ^a
6	2.20±0.44	4.01±1.00 ^c
7		4.15±0.78 ^b
8	4.58±2.18	5.41±0.78 ^b
11		7.27±2.67

Values significantly different from the first value of the same column are marked with *a*: (P<0.05), *b*: (P<0.005) or *c*: (P<0.001).

TABLE III

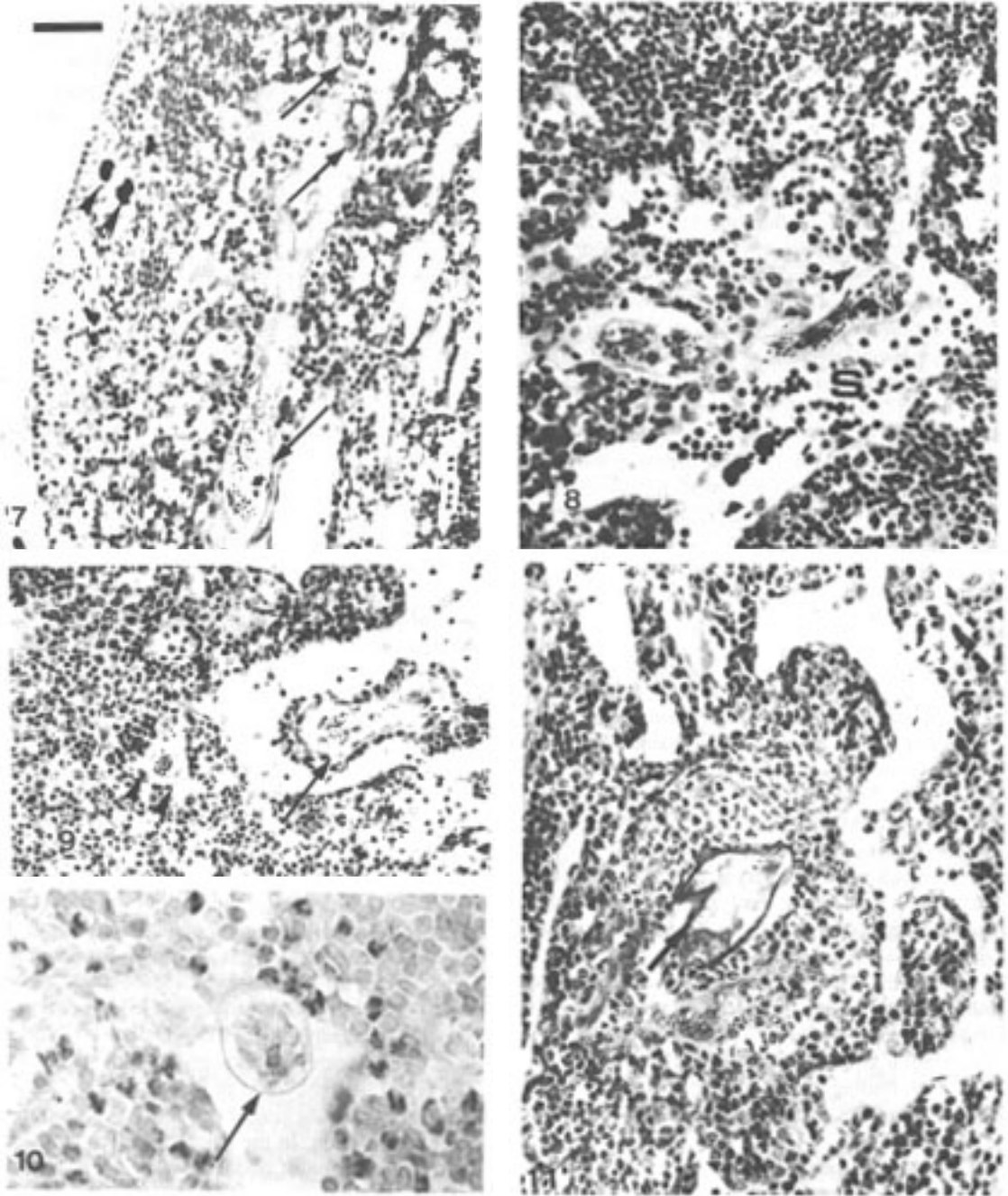
Estimative of the amount of schistosomules in the right parotid lymph nodes in naive and infected (immune) Balb/c mice. Infection or re-infection with *Schistosoma mansoni* cercariae through the right ear. The number of animals in parenthesis

Days after (re)infection	Naive mice	Re-infected mice
1		no larva (6)
2	no larva (4)	no larva (6)
3		0-3 larva/section of 2 nodes (5)
4	one larva in one node (5)	1-5 larvae/section, all nodes (4)
5		1-7 larvae/section, all nodes (4)
6	1-3 larvae/section, all nodes (5)	0-4 larvae/section, all nodes (5)
7		1 larva in 1 node (4)
8	0-2 larvae/section of 3 nodes (5)	no larva (5)
11		no larva (4)

ected animals, the larvae were practically absent till day 4 of infection, but they were found in all lymph nodes at day 6, the number decreasing by day 8. In the challenged mice, a few larvae appeared by day 3 of re-infection, all lymph nodes exhibiting schistosomula from day 4 to 6 of re-infection. The larvae became rare at days 7 and 8 and had disappeared by day 11. The larval distribution inside the lymph nodes was the same at infection and re-infection. Most larvae lay in the subcapsular sinus (Fig. 3) or in the perihilar segments of medullary sinuses (Fig. 7). A few larvae were seen in extranodular zones and in the segments of the medullary sinuses just below these zones, but they were excluded from the deep cortical units and center of lymphoid nodules. The larger number of larval profiles in the re-infected mice allowed some additional information on the

larval migration. At least one larva was found in a small efferent lymph vessel. Serial sections demonstrated that the same larva could have a part in a medullary sinus and another in a medullary cord (Fig. 8), sometimes adjacent to its vein. Finally, two schistosomula were seen in the lumen of medullary cord veins at day 5 of re-infection (Fig. 9).

At day 6 of the primary infection, all nodes exhibited eosinophilic inflammatory processes in some cortical extranodular zones, lying between two deep cortex units or in contiguity with the perihilar region (Fig. 4). Some medullary cords, close to the extranodular zones or deep cortex units, also exhibited a higher concentration of eosinophils. At the periphery of deep cortex units, the high endothelial venules showed several eosinophils in their lumen, wall, and neighboring extranodular tissue (Fig. 5). The medullary cord



Parotid lymph nodes of infected mice killed at days 5-7 of re-infection with cercariae of *Schistosoma mansoni*. Toluidine blue-borate, excepting Fig. 10. Fig. 7: schistosomulum profiles in the perihilar region (arrows). Fig. 8: a larva passing from a medullary sinus to a medullary cord. Fig. 9: a medullary cord vein presenting a larva in its lumen (arrows). Fig. 10: a larva (arrow) in a medullary sinus exhibiting eosinophils adhered to its surface. Giemsa stain. Fig. 11: medullary region showing a degenerating larva (arrow) surrounded by eosinophil nuclei. Bar = 48 μ m (7.9); 40 μ m (8); 19 μ m (10); 30 μ m (11).

veins close to the deep cortex units could also exhibit eosinophils in their lumen and wall (Fig. 6). No schistosomulum was seen inside the eosinophilic infiltrates and all larvae were free of adher-

ent granulocytes, looking intact. At day 8, eosinophils were frequent in medullary cords, being rarely found in the lymphatic sinuses, however, none contacted the larvae that remained intact.

In contrast to the findings in primarily-infected animals, a few larvae exhibiting clear signs of degeneration were observed at days 5, 6 and 7 of re-infection. Inflammatory processes were absent or very discrete till day 4 of re-infection. The presence of eosinophils in the lumen and wall of medullary cord veins was frequent, mainly at day 7. At day 5, only one lymph node presented a focal and intense eosinophilic inflammatory process located in the medullary region and around a degenerating larva. However, eosinophils were easily found in medullary cords of all lymph nodes as well as in some extranodular zones. Some larvae exhibited eosinophils contacting their surface (Fig. 10). The eosinophilic infiltrates became more frequent at days 6 and 7 of re-infection, most of them being discrete and keeping the distribution already described. Intense eosinophilic processes were restricted to the extranodular zone (just one), or medullary cords (two nodes). Each one of these two medullary cords infiltrates surrounded a degenerating larva (Fig. 11). The eosinophilic infiltrates became very discrete at day 8, being absent at day 11 of re-infection.

DISCUSSION

The parotid lymph node of the mouse exhibited the overall architecture described for the non-segmented lymph nodes of rats (Sainte-Marie et al. 1982), even regarding the distribution of mast cells (Sainte-Marie & Peng 1990). Our findings showed that schistosomula can pass through regional lymph nodes without being damaged, even in challenged mice. The larval migration route coincides with the flow of the lymph, as already described in naive mice (reviewed by Wheater & Wilson 1979). However, we have succeeded in demonstrating that the larvae can also leave the node by penetrating the veins of the medullary cords, probably after migration through extranodular zones in contact with medullary sinuses. This finding indicates that schistosomula can guide themselves to veins, even when in a lymphatic channel.

Contrasting with the skin reaction induced by cercarial penetration (Incani & McLaren 1984), the lymph nodes of either infected or re-infected mice presented rare neutrophils. Studies on the peritoneal cavity of naive mice have shown that cercarial inoculation provokes an influx of neutrophils that adhere to the cercarial surface and remove fragments of glycocalyx (Melo et al. 1993a, b). Our results indicate that when schistosomula reach the nodes, they are probably devoid of molecules able to attract neutrophils or mediate their adhesion to the larval surface.

In rodents, the number of mast cells increases

rapidly in regional lymph nodes after antigenic or non-immunogenic stimulation (Miller & Cole 1968, Roberts 1970, Wlodarski et al. 1973), probably by a process of draining from the stimulation site. However, within 24-48 hr, the number of nodal mast cells decreases to values below the normal ones (Wlodarski et al. 1973). In our primarily-infected mice, the mast cell number increased till day 4 after stimulation dropping to control values only at day 8. Probably, these alterations followed the phenomena occurring at the site of cercarial stimulation. After cercarial penetration, the skin thickness attains maximal values in 48 hr (Incani & McLaren 1984), and this is consistent with the rise in mast cell number at day 2 with peak at day 4. Also, by remaining in the skin for 4 to 5 days (Miller & Wilson 1978), or up to 7 days (Gerken et al. 1986), the schistosomula could create conditions able to maintain the raised number of nodal mast cells till day 6.

In the absence of challenge infection, the parotid lymph nodes of infected mice exhibited a larger number of mast cells in comparison with normal animals. Accordingly, this number is also enlarged in the dermis (Fernex & Fernex 1982), peritoneal cavity and liver granulomas (Lenzi et al. 1987), and the histamine levels are higher in the ears (Gerken et al. 1990). The re-infection induced a rise in nodal mast cell number, but there was a delay in comparison with the primary infection. This delay may involve differences in mast cell degranulation.

In the primary infection, the number of degranulating mast cells was significantly higher at day 4, occurring after the peak of the skin reactions. In contrast, the number of degranulating mast cells increased at the first day of re-infection as expected for IgE bearing mice (Mota-Santos et al. 1985). In the skin of re-infected mice, the reaction also involves early mast cell degranulation (Bentley et al. 1981, Incani & McLaren 1984). Both dermal and nodal degranulation processes could account for the delay in the rise of nodal mast cell number after re-infection. In the primarily-infected mice, the degranulation of nodal mast cells was not related to the presence of larvae or inflammatory process. Since IgE induced by larval antigens is not supposed to be present at day 4 of infection, direct effects of larval antigens or cross-reactions could be involved.

The richness in eosinophils at sites of parasite infection has been recognized for many years. However, the role played by these cells is not completely elucidated. *In vitro* studies have shown that eosinophils might be efficient anti-parasite effectors by their ability to kill schistosomula in the presence of immune sera (Butterworth et al. 1975, Sher

& Rees 1975, Anwar et al. 1979). However, *in vivo* evidence for such a role is sparse and equivocal (Mahmoud et al. 1975, Sher et al. 1990). In the present study, both primary infection and re-infection induced an influx of eosinophils to nodal tissue through the high endothelial venules and their collecting veins in medullary cords. However, only in the re-infected animals were the eosinophils able to adhere to the larval surface, damaged larvae being found inside eosinophilic infiltrates. The significance of this adhesion is not clarified by our histological study, since it could either signify a role in larval death or a phenomenon secondary to larval degenerative changes. Notwithstanding, the regional lymph nodes seem to be of slight importance in limiting the re-infection, judging by the small number of intranodal degenerating larvae. However, larval lesions could have occurred without detectable histologic changes. Therefore, it remains the possibility of larval damage inside and death outside the nodal tissue.

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