INTRODUCTION

Kinetoplastid flagellates occur as extracellular blood parasites in the vascular system of a wide variety of different fishes. In European cyprinids, the trypanosomatid *Trypanosoma carassii* (formerly known as *T. danilewskyi*) and the bodonid *Trypanoplasma borreli* are widely distributed (Lom 1979, Lom & Dykova 1992). They are transmitted by leeches, which leads to an initial rise in blood parasitaemia, followed by a decline in parasite numbers. Thereafter, low numbers of flagellates are present in the blood and internal organs of most fishes for a prolonged period. Chronic infections may result in a high prevalence of the flagellates in a fish population, but the intensity of infection is generally low (Lom 1979, Steinhagen et al. 1989, Lom & Dykova 1992, Jones et al. 1993).

Infections with *Trypanosoma carassii* and *Trypanoplasma borreli* result in a characteristic non-fluctuating parasitaemia (Steinhagen et al. 1989, Overath et al. 1999). Fish that have controlled the acute infection are resistant against infection with lines of *T. carassii* or *T. borreli*, isolated from other carp in the chronic phase of infection (Overath et al. 1999, D.S. unpubl. obs.). Parasite-specific antibodies are believed to control these infections. They can be detected in the sera of
infected fish (Lom & Dykova 1992, Jones et al. 1993, Wiegertjes et al. 1995, 1996), and sera (or immunoglobulin, IgM) of convalescent fish can transfer resistance to infection with *T. carassii* (Overath et al. 1999) to naïve carp or can reduce the parasitaemia after a *T. borreli* infection (Wiegertjes et al. 1995). Complement fixing antibodies have also been observed to protect rainbow trout *Oncorhynchus mykiss* (Li & Woo 1995, Feng & Woo 1997) and *Salvelinus fontinalis* (Ardelli & Woo 1997) from infection with *Cryptobia salmositica*, a haemoflagellate closely related to *T. borreli*. Plasma from naïve *S. namaycush*, naturally resistant to *C. salmositica*, lysed the parasite in vitro, indicating that alternative complement activation might be the mechanism of resistance (Ardelli & Woo 1997).

Some strains of carp were found to be highly susceptible to infection with *Trypanoplasma borreli*. Infection usually results in a 100% mortality 3 to 4 wk after injection of the flagellates. In the serum of these fish, parasite-specific antibodies could not be detected (Wiegertjes et al. 1995). In addition, these *T. borreli*-susceptible carp did not mount an antibody response to an unrelated antigen (dinitrophenyl-keyhole limpet haemocyanin, DNP-KLH). Thus it was considered that these carp might have a genetically predetermined low antibody response (Wiegertjes et al. 1995).

In mammalian trypanosomiasis, besides an antibody response, the state of macrophage activation is considered to be critical to trypanotolerance (Tabel et al. 2000, De Baetselier et al. 2001). Salivarian trypanosomes were able to induce a secretion of inflammatory molecules such as NO (nitric oxide), TNF (tumor necrosis factor) or IL1 (interleukin) by activated murine macrophages, which inhibited T-cell proliferative responses to parasite-related and -unrelated antigens (Schleifer & Mansfield 1993, Sternberg 1998). In addition, De Baetselier et al. (2001) found that certain *Trypanosoma brucei* strains induced different, alternative cytokine patterns, which correlated with their pathogenicity.

In order to further characterise the role of a specific immune response for defence against *Trypanoplasma borreli*, we analysed serum from susceptible and resistant carp (collected after *T. borreli* challenge) for the ability to kill *T. borreli* and for the presence of *T. borreli*-specific immunoglobulin. In order to determine whether in the context of a *T. borreli* infection leukocyte-derived molecules suppress lymphocyte activation and thus might contribute to increased parasite susceptibility, we investigated the ability of *T. borreli*-stimulated head kidney leukocytes (HKL) and peripheral blood leukocytes (PBL) from carp strains of different parasite susceptibility to modulate lymphocyte activation in response to mitogens (pokeweed mitogen, PWM, and phytohaemagglutinin, PHA).

**MATERIALS AND METHODS**

*Trypanoplasma borreli*-susceptible carp and resistant pond carp. Susceptible carp: Carp of a single crossing (E20 × R8, Wageningen Agricultural University, The Netherlands), susceptible to *T. borreli* infection, were bred and raised at 20 to 23°C in recirculated filtered tap water. After initial feeding with *Artemia* sp. nauplii for 4 wk, the food was switched to pelleted dry food (Milkivit). Carp, 2 yr old and weighing 200 to 300 g, were used for blood collection and infection with *T. borreli*. Before infection, the carp were acclimatised to a re-circulating system of separate 120 l tanks at 20 ± 1°C for at least 2 wk.

Resistant carp: We obtained 2 yr old carp (body weight 200 to 300 g) from a hatchery in the vicinity of Hannover (Germany). The carp were selected from pond carp and transported to the laboratory, where groups of 20 fish were maintained in recirculated, filtered tap water at room temperature (18 to 20°C) in 300 l tanks and fed daily with pelleted dry food (Milkivit). Carp were adapted to the maintenance conditions for at least 8 wk before infection or leukocyte sampling. *Trypanoplasma borreli* flagellates in the circulating blood were never observed (checked microscopically at least 4 times), a previous infection with the blood flagellate, however, could not be excluded.

Infection experiments. *Trypanoplasma borreli* had previously been cloned and characterised, and was maintained in the laboratory by syringe passage as described earlier (Steinhagen et al. 1989). To check the carp for susceptibility to the *T. borreli* clone, 12 carp from each group were each injected intramuscularly (i.m.) with $1 \times 10^4$ *T. borreli* in 100 µl phosphate buffered saline (PBS). Every 2 wk (up to 8 wk post-infection), all carp were examined for the presence of *T. borreli* flagellates in the circulation: the fish were anaesthetised in 0.15 g l$^{-1}$ tricaine methane sulphonate (MS 222, Sigma-Aldrich); blood was collected from each carp by gill puncture with a glass capillary, transferred to glass slides and microscopically monitored for the presence of parasites.

For serum collection from infected carp, 3 resistant and 3 susceptible carp were injected intramuscularly with $1 \times 10^4$ *Trypanoplasma borreli* in 100 µl PBS. Control carp (3 carp from both groups) were injected with 100 µl PBS. Infected and control carp were kept under identical conditions. At Day 14 post-injection (p.i.), all carp were killed by immersion in 0.5 g l$^{-1}$ MS 222. Whole blood was collected from the caudal vein and transferred to polystyrene tubes. The blood was kept at 22°C for 4 h followed by 48 h at 4°C. After centrifugation (15 min at 750 × g), the serum was collected and the sera within the groups were pooled and kept frozen at −80°C until use. For each fish, the para-
Sitaemia of *T. borreli* was determined by counting the flagellates present in the blood samples, using a Neubauer counting chamber.

**Separation and blotting of carp serum proteins.** Serum proteins were separated under reducing conditions by discontinuous sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) as described by Wagner et al. (1995). Sera were diluted 1:50 in sample buffer and 10 µl were run per lane. Gels were either stained for proteins with Coomassie blue according to standard protocols (Sambrook et al. 1989) or blotted on PVDF-membranes (Millipore). Blots were incubated with the murine monoclonal antibody WCI 12 (specific for carp heavy-chain IgM; Secombes et al. 1983), and subsequently with a polyclonal goat anti mouse IgG (H+L) conjugated with alkaline phosphatase (Dianova). The substrate reaction was based on the method described by Blake et al. (1984).

**Culture media.** Culture media for cell separation and cultivation as well as PBS were diluted with distilled water (10% v/v) to adjust their osmotic pressure to carp serum osmolarity. Diluted RPMI 1640 (Biochrom) with 50 000 IU l⁻¹ sodium heparin (Sigma-Aldrich) was used for blood collection (heparinised medium). Diluted RPMI 1640 with 10 000 IU l⁻¹ sodium heparin served as washing solution. For cultivation experiments, diluted RPMI was supplemented with 1% (v/v) ‘carp serum’ (leukocyte culture medium). Carp serum was a pool of heat-treated (30 min, 56°C) and sterile-filtered (0.2 µm) sera from 15 individual fish.

*Trypanoplasma borreli* were raised in a mixture of Hanks buffered salt solution (42.5%, v/v), Earl’s modified minimum essential medium (21.25%, v/v), Leibovitz 15 medium (21.25%, v/v), distilled water (10%, v/v) and 5% carp serum (HML medium, Steinhagen et al. 2000). All culture media were supplemented with 5% CO₂. All set-ups were in triplicate. PBL were stimulated with pokeweed mitogen (PWM; 1 mg l⁻¹), phytohaemagglutinin (PHA; 3 mg l⁻¹), and viable or lysed *Trypanoplasma borreli* (5 × 10⁸ cells ml⁻¹).

For the preparation of culture supernatants (SN) PBL and HKL (5 × 10⁶ ml⁻¹) from susceptible and resistant carp were incubated in 24-well flat-bottomed plates in a final volume of 1 ml well⁻¹ for 4 d. Parallel set-ups contained cells together with 2.5 × 10⁶ viable *Trypanoplasma borreli* ml⁻¹ or 2.5 × 10⁶ viable *T. borreli* ml⁻¹ alone. All assays were made with cells of 3 resistant and 3 susceptible carp. The culture supernatants of individual set-ups were pooled, centrifuged (15 min, 750 × g) and stored at −80°C until use.

**Leukocyte cultivation.** For proliferation experiments, peripheral blood leukocytes were incubated in 96-well flat bottomed microtitre plates (10⁶ cells well⁻¹) in a final volume of 175 µl leukocyte culture medium for 4 d at 27°C in water vapour-saturated atmosphere with 3% CO₂. All set-ups were in triplicate. PBL were stimulated with pokeweed mitogen (PWM; 1 mg l⁻¹), phytohaemagglutinin (PHA; 3 mg l⁻¹), and viable or lysed *Trypanoplasma borreli* (5 × 10⁸ cells ml⁻¹).

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**Flow cytometry analysis of cultured cells.** Suspensions of carp leukocytes (PBL, HKL) and *Trypanoplasma borreli* were analysed with a FACScan® (Becton Dickinson, single excitation wavelength of 488 nm). Cellular subsets were identified according to their characteristic forward and side-scatter values (FSC/SSC profiles; Verburg-Van Kemenade et al. 1994, Scharsack et al. 2000). Propidium iodide (2 mg l⁻¹ final conc.) was added to the suspensions to label membrane-damaged cells. At least 10 000 events were acquired and analysed with the software WinMDI,
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Version 2.8 (Trotter 1998: WinMDI 2.8 software package; www.facs.scripps.edu/software.html). Cellular debris with low FSC characteristics and dead cells were excluded from further evaluation.

Total cell numbers were determined with the single-cell dilution assay (Pechhold et al. 1994) using bovine, FITC-labelled PBL as standard cells (Hendricks et al. 2000). Numbers (N) of cultivated cells in individual wells were calculated according to: 

\[ N_{\text{viable cells}} = \frac{\text{events}_{\text{viable cells}} \times N_{\text{standard cells}}}{\text{events}_{\text{standard cells}}} \]

Statistics. Data sets were compared by ANOVA and Duncan’s multiple-range tests. Differences with \( p < 0.05 \) were considered significant.

RESULTS

Infection experiments

All 12 susceptible carp showed parasitaemia and died between 21 to 25 d after infection with Trypanoplasma borreli flagellates. In contrast, \( T. borreli \)-injected pond carp (n = 12) did not show any sign of parasitaemia throughout the observation period of 8 wk.

Sera were taken at Day 14 post-infection. The pooled sera of both susceptible and resistant carp appeared to contain some immunoglobulins, and in both groups infection with \( T. borreli \) did not result in a change in the total serum immunoglobulin concentration (Fig. 1, cf. Lane 1 with Lane 2 or Lane 3 with 4).

Striking differences appeared when the sera were tested for \( T. borreli \)-specific antibodies by incubation with live \( T. borreli \) flagellates. Sera from PBS-injected, resistant carp already contained \( T. borreli \)-reactive antibodies (Fig. 2), which may indicate previous infection with the parasite. The higher fluorescence intensity at lower serum dilution and the higher fraction of stained \( T. borreli \) flagellates (Fig. 2) indicates that specific antibodies were boosted after infection of resistant carp with \( T. borreli \). In contrast, no reactivity of serum antibodies from PBS-injected or \( T. borreli \)-infected susceptible carp with the flagellates was detectable (Fig. 2).

Incubation of \( T. borreli \) with serum from challenged as well as non-challenged pond carp killed the parasites after 18 h \textit{in vitro} (Table 1). This trypanocidal activity of the sera disappeared after heat treatment of the sera. This could indicate that complement activity is involved in the killing of trypanoplasms \textit{in vivo}. The viability of \( T. borreli \) was not affected by sera of susceptible carp (Table 1).

In \textit{vitro} responses of head kidney leukocytes and peripheral blood leukocytes from carp resistant and susceptible to \( T. borreli \)

The inability of susceptible carp to produce antibodies after infection with \( T. borreli \) was not due to a general inability of peripheral blood leukocytes to respond to mitogens (PWM, PHA) with proliferation (Fig. 3). Also the blastogenesis in response to
lysed T. borreli preparations was comparable between PBL of resistant and susceptible carp (Fig. 3). However, differences were apparent when PBL were cultured in the presence of viable flagellates: PBL from resistant carp responded with blastogenesis in a magnitude comparable with PHA-stimulated PBL, whereas the response of PBL from susceptible carp towards viable T. borreli was significantly lower (Fig. 3).

A Trypanoplasma borreli-induced proliferative response of PBL from resistant carp was also observed with cell-free culture supernatants (SNs) of the flagellates (Fig. 4, Set-up 2). Again, the responses of PBL from susceptible carp were not modulated by SNs from T. borreli (Fig. 4, Set-up 1).

Differences in the responsiveness of PBL from both carp groups could be due to a different range of secreted mediators. We therefore cultivated PBL of resistant and susceptible carp and tested the cell-free supernatants for their modulatory potential in stimulation assays.

While SNs of PBL from resistant carp clearly enhanced the PWM- and PHA-induced blastogenesis of PBL from resistant carp, this effect was significantly less apparent with PBL from susceptible carp (cf. Set-ups 3 and 4, Fig. 4). Interestingly, SNs from cultured PBL of susceptible carp enhanced the proliferative response of PBL from susceptible carp, however, the PHA-induced response of PBL of resistant carp was only slightly enhanced, and the PWM-induced response was even diminished (cf. Set-ups 5 and 6, Fig. 4).

The differential modulatory capacities of SNs from cultured PBL indicate that the cells of resistant and susceptible carp produce a different range of soluble mediators.

This was further substantiated when SNs which were produced by PBL in the presence of Trypanoplasma borreli were tested. SNs of resistant carp stimulated or enhanced the proliferative response of PBL from resistant carp, but not that of PBL from susceptible carp (cf. Set-ups 7 and 8, Fig. 4). Interestingly, SNs from PBL of susceptible carp increased the proliferative response of both PBL from resistant carp and that of PBL from susceptible carp (cf. Set-ups 9 and 10, Fig. 4).

In contrast to PBL, HKL of susceptible and resistant carp did not differ much in their spectrum of secreted mediators.
and modulating mediators: HKL supernatants of both carp lines suppressed the mitogen-induced proliferation of PBL from susceptible carp (especially the PWM-induced proliferation; data not shown). Also, when HKL were cultivated in the presence of *Trypanoplasma borreli*, the supernatants of cells of both carp lines suppressed the cellular proliferation of PBL from resistant or susceptible carp largely to the same degree (data not shown).

**DISCUSSION**

Carp susceptible to an infection with *Trypanoplasma borreli* died after experimental infection. Sera from this strain of carp contained no detectable antibodies, either before or after infection. Despite the limited number of individuals analysed here, these results support previous findings (Jones et al. 1993, Wiegertjes et al. 1995), which suggested that generation of a specific immune response is decisive for effective elimination of *T. borreli*.

In contrast, the pond-reared carp contained antibodies both before and after experimental infection. Heat-labile factors, most probably complement, were responsible for effectively killing the parasite in vitro. As the parasite was killed in the presence of antibodies binding to *Trypanoplasma borreli*, complement activation by the classical pathway could be suspected. Complement activity also was observed to be important in the protection of salmonid fish from infections with *Cryptobia salmositica*. In rainbow trout infected with *C. salmositica*, complement-fixing antibodies were shown to be the key mechanisms combating the infection (Li & Woo 1995, Feng & Woo 1997), while in *Salvelinus namaycush*, a fish species with natural resistance to *C. salmositica* complement activated via the alternative pathway mediated the destruction of the parasites (Ardelli & Woo 1997). Our experiments cannot exclude a possible role of alternatively activated complement in resistance of carp to *T. borreli* infection, but the complete lack of trypanocidal activity in serum from susceptible carp suggests that in serum from these carp, complement activation was not via the alternative pathway in the presence of *T. borreli*.

The 2 different carp strains investigated here did not only differ in genetic background, but also in life history. While carp susceptible to the flagellate were...
grown under laboratory conditions, the resistant carp were obtained from a local hatchery. Thus differences of the fish groups in their response to *Trypanoplasma borreli* besides their different genetic background might be influenced by their different life histories as well.

The reason for the lack of antibody response in susceptible carp is still unclear. Serum samples for the present investigation were collected on Day 14 p.i. Upon injection of sheep red blood cells (SRBC), carp showed a plaque-forming cell response which peaked on Day 12 p.i. when the fish were kept at 20°C, and on Day 9 at 24°C (Rijkers et al. 1980). In the present study, carp were kept at 20°C; thus the presence of some antibodies might be expected in their serum by 14 d post-infection with the parasite. However, Wiegertjes et al. (1995) could not detect antibodies to *Trypanoplasma borreli* or to an antigen unrelated to the parasite (DNP-KLH) in the serum of individuals from the same carp line even 21 d p.i. Therefore, Wiegertjes et al. (1995) suggested a more general B-cell anergy in these carp. However, when peripheral blood lymphocytes (PBL) were stimulated in vitro, cells of both resistant and susceptible carp responded with blastogenesis after mitogenic stimulation in a comparable manner. Previously, we also showed in blood, head kidney and spleen of *T. borreli*-infected susceptible carp the presence of many lymphoblasts, among which surface immunoglobulin-positive B-cells occurred (Barckhausen-Kiesecker 1995, Scharsack et al. 2000). In addition, sera from susceptible carp contained detectable amounts of heavy-chain Ig, indicating a general ability of susceptible carp to produce Ig.

Although we did not check in this study whether B-cells were activated or not, a *Trypanoplasma borreli*-induced proliferation of PBL could be observed in susceptible carp (Scharsack et al. 2000, 2003a). However, cells of susceptible carp proliferated significantly less in response to viable *T. borreli* compared to cells from resistant carp. This could be due to a memory effect in resistant carp resulting from a previous pond infection with *T. borreli*. In serum from resistant carp, considerable amounts of *T. borreli*-specific antibodies were detected, even before infection under laboratory conditions. Nevertheless, infection with *T. borreli* resulted in a boosting of Ig production in resistant carp, while in infected susceptible carp no *T. borreli*-specific Ig was detectable.

If B-cells of susceptible carp are not anergic and are stimulated by *Trypanoplasma borreli* in vivo (Barckhausen-Kiesecker 1995) and in vitro, the inability of susceptible carp to mount an Ig response could be related to an insufficient modulatory impact.

One possible explanation could be a lack of an appropriate mediator production by accessory mono-
soma cruzi infection in man, abrogation or reduction of a strong polyclonal lymphocyte activation correlates with an increased resistance to infection (cf. Minoprio 2001). In a similar way, the differences between the carp lines in susceptibility to T. borreli could be based on a strong polyclonal activation, which is not down-regulated by appropriate inhibitory signals provided by yet unidentified cellular lymphoid subsets.

LITERATURE CITED


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