The effects of tumour necrosis factor-α and interleukin1 on an in vitro model of thyroid-associated ophthalmopathy; contrasting effects on adipogenesis

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Introduction

Thyroid-associated ophthalmopathy (TAO) is an autoimmune inflammatory condition that is also known as thyroid eye disease and Graves’ ophthalmopathy. TAO affects between 25 and 50% of those with autoimmune hyperthyroidism, with 3–5% suffering severe eye disease, which may threaten vision (1, 2). The pathogenesis is not fully understood, but the key pathological processes include acute inflammation, excess production of glycosaminoglycans (GAG) and expansion of adipose tissues within the orbit (3). The treatment for more severe cases of TAO is often unsatisfactory and there has been increasing interest in the possibility of using anti-cytokine agents to treat active TAO (4, 5). This idea is based on in vitro data on the effects of cytokines on orbital fibroblasts (OFs) and the growing experience of the use of anti-cytokine agents in other inflammatory conditions, such as rheumatoid arthritis and inflammatory bowel disease. Cytokines, such as interferon γ (IFNγ), interleukin 1 (IL1) and tumour necrosis factor-α (TNFα) have been shown to stimulate GAG production from OFs in vitro (6–9). These same cytokines have been shown to increase ICAM1 expression using immunohistochemical techniques (10) and are also increased in active TAO in vivo (11).

These and other data suggest that blocking these cytokines might inhibit some of the pathogenic processes in TAO, including the orbital inflammatory response and excess GAG production, therefore limiting orbital tissue expansion. However, there are data that show that TNFα can inhibit orbital adipogenesis, and hence blocking TNFα might have a harmful effect on TAO by increasing adipogenesis and increasing orbital tissue volume (12).

We aimed to evaluate the effects of selected cytokines and anti-cytokine agents on cultured OFs as an in vitro model of TAO. We focused on TNFα and IL1, because anti-cytokine agents that block the action of these cytokines are presently in clinical use for indications other than TAO. We chose to examine three endpoints: ICAM1 expression, GAG production and adipogenesis. Increased ICAM1 expression plays a role in inflammatory cell
recruitment and occurs in active TAO (11). Thus, a reduction in ICAM1 expression would be expected to have an inhibitory effect on the local inflammatory response. We measured GAG production by orbital fibroblasts and adipogenesis in pre-adipocyte orbital fibroblasts, as these processes are thought to cause tissue swelling in TAO (1, 13).

Subjects

TAO was diagnosed on clinical grounds based on the presence of typical clinical features in the context of autoimmune thyroid disease. In the TAO group, seven out of the eight patients were female, median age 49 years (range 42–65), median Mourits activity score was 1 (range 0–9), and five of the eight patients were smokers (Table 1). Control subjects were patients without a personal or family history of thyroid disease or TAO, and without clinical evidence of the same, who were attending orbital surgery. In the control group, four out of the five patients were female, median age 48 years (range 21–77) and all were non-smokers (Table 1). The study was approved by the institutional ethics committees and all study participants gave written informed consent.

Methods

Orbital fibroblast cultures

Orbital fibroblast (OF) cultures were established, similar to published methods (14). Orbital fat biopsies (approximately 4–8 mm in length) were taken at surgery and transported to the laboratory in normal saline at room temperature. The biopsies were diced using a scalpel blade and placed in RPMI tissue culture medium containing 9% fetal bovine serum, 20 ml/l of 1 M HEPES, 1.8 mg/l fungizone, 35 U/ml penicillin and 35 μg/ml streptomycin in a 25 cm² flask and grown at 37°C in 5% CO₂ and 21% O₂ (all additives from Gibco, Invitrogen). Once OFs were adherent, the flask was washed with culture media and OFs grown to confluence replacing media every 3–4 days. Confluent fibroblasts were passaged and plated into new flasks or multi-well plates, with typically 20 000 cells being seeded into each well of a 12-well plate, which could be grown to confluence in 5–10 days.

Cytokines (TNFα, IL1, IFNγ, transforming growth factor-β (TGFβ) and IL10) were purchased from R&D Systems and the concentrations chosen from logarithmic dose–response experiments, they being on either side of the ED50. The concentrations of anti-cytokine agents used were based upon published levels of peak serum concentrations in humans following typical therapeutic doses (anti-TNFα agents adalimumab 50 μg/ml (15), infliximab 150 μg/ml (16) and etanercept 2.5 μg/ml (17) and anti-IL1 agent anakinra 5 μg/ml (18)), and a concentration 1/100 of the same.

ICAM1 measurements

The expression of ICAM1 by OFs was analyzed using flow cytometry. OFs were grown to confluence in 12-well plates. Confluent OFs were given fresh medium and stimulated for 24 or 48 h with the cytokine ± anti-cytokine agent, with un-stimulated wells as control. OFs were trypsinized and centrifuged, and resuspended in 200 μl of culture medium. OFs were then added to 5 μl of

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n/a, not applicable.

*Clinical activity score refers to the Mourits clinical activity score (37), with 10 representing the most active disease and 0 the least active disease.
isotype IgG control PE mouse antibody (BD, Ref 345816) or 5 μl of anti-ICAM1 PE antibody (BD, Ref 555511), vortexed and incubated in darkness at 4 °C for 30 min. OFs were washed with phosphate buffer and resuspended in 1% paraformaldehyde. The OFs were analyzed using a BD FACScan cytometer and Cellquest analysis software with data expressed as median fluorescent intensity of each experimental condition. Inter- and intra-assay coefficient of variation (CV) values were 2.7 and 2.2%, respectively.

**GAG measurements**

The level of hyaluronic acid (HA) in the supernatants from the 48-h stimulation experiments described earlier was measured using an ELISA kit (K-1200, Echelon Biosciences, Inc., Salt Lake City, UT, USA) as per the manufacturer’s instructions. Samples were diluted 1/10 before analysis, and the average of duplicate measurements was taken. Inter- and intra-assay CV values were 10.8 and 6.6%, respectively.

**Adipogenesis**

Orbital fibroblasts were exposed to a differentiation protocol to encourage adipogenesis according to published methods (12, 19–22). OFs were grown to confluence in 6- or 12-well plates and exposed to differentiation media for 4 days (RPMI as detailed earlier, but lacking fetal bovine serum, and supplemented with 0.1 mM IBMX, 1 μM dexamethasone, 1 μM rosiglitazone (Avandia, GSK), 100 nM insulin (Actrapid, Novo Nordisk), 10 μg/ml transferrin (Sigma T 3309), 0.2 nM T3 triiodothyronine (Fluka 91 990). Media was replaced at days 4 and 8 with propagation media (identical to differentiation media but lacking the IBMX, dexamethasone and rosiglitazone) and cells analyzed at day 10. Cytokines and anti-cytokine agents were added at day 1, and continued for the 10 day differentiation protocol, being replaced whenever media was replaced. Cells were stained for lipid using oil-red-O (ORO), photographed, and assessed for degree of adipogenesis using three methods: (i) extraction of ORO and measuring absorbance units (AU) at 492 nm (23), (ii) a visual scoring system (Table 2), and (iii) computer analysis of digital photographs.

### ORO staining and adipogenesis quantification

ORO (0.7 g, Sigma O 0625) was dissolved in 200 ml isopropanol (Sigma I 9030), shaken and left overnight at 21 °C. The solution was repeatedly centrifuged at 13 000 r.p.m. and decanted, until no un-dissolved debris remained. This solution was diluted 3:2 with distilled water and left overnight at 4 °C. This 60% ORO solution was repeatedly centrifuged and decanted until no un-dissolved debris remained, and was then ready for use.
OFs ready for analysis were washed twice with PBS before being rinsed in 60% isopropanol for 30 s. 200 μl 60% ORO solution was added for 10 min, before being removed and each well rinsed with 200 μl 60% isopropanol for 10 s. OFs were rinsed twice with PBS and photographed with a digital camera. The lipid droplets in the cells were stained orange-red and visually assessed for degree of adipogenesis. The wells were then washed four times with PBS to remove any remaining un-bound ORO. PBS was removed and 200 μl 100% isopropanol was added for 5 min to extract the ORO. A volume of the supernatant (170 μl) was transferred to a reading plate and absorbance units measured at 492 nm. Since no individual well can be measured more than once, it was not possible to calculate inter- and intra-assay CV values for this method. However, in measuring wells containing cells from different patients, but under the same experimental conditions, the inter- and intra-assay variations (which includes inter-sample variation) were 26.9 and 8.5% respectively.

In order to assess adipogenesis according to the microscopy appearance of the OFs, a visual scoring system was designed (Table 2). This score was based upon the number of high-power (200×) microscopy fields containing cells with lipid droplets, the number of cells per field containing lipid droplets and the average size of the lipid droplet (in the present study this multiplication factor for droplet size was not used as droplet size was relatively uniform). Approximately, ten fields in each well were examined, though more were viewed if there was uncertainty. The inter- and intra-assay CV values were 14.3 and 8.3% respectively. In addition, a microscopy analysis software package (AnalySIS, version 3.2, Soft Imaging Systems GmbH, Münster, Germany) was used to assess photographs of OFs stained with ORO taken by a digital camera mounted on a microscope. The software quantified the percentage of pixels in the photograph with a set degree of red, blue and green that corresponded to the areas of ORO staining. The inter- and intra-assay CV values were 21.3 and 1.7% respectively.

Statistics

The paired t-test was used to test ICAM, GAG and adipogenesis responses compared with the un-stimulated control condition from each experiment. Correlation of different methods of assessing adipogenesis was done using Pearson product moment correlation. Statistical significance was taken to be <0.05 and data analyzed using Minitab 13 software.

Results

ICAM1 measurements; cytokine responses

Experiments using different cell-detachment techniques (cell scraper, trypsin (0.25%), trypsin (0.25%)/EDTA (1 mM) and EDTA (5 mM)) prior to flow cytometry analysis found that the relative ICAM1 expression was similar across all methods, although the absolute measurements were 25–30% greater with trypsin/EDTA and EDTA. Trypsin (0.25%) was used for this study. Dose–response curves were performed for each cytokine and where a dose–response was seen, concentrations either side of the ED50 were used on OFs from five patients with TAO and five control subjects. TNFα, IL1 and IFNγ stimulated ICAM1 expression in OFs from patients with TAO and control subjects in a similar, dose-dependent manner. TNFα (0.1 ng/ml), IL1 (0.1 ng/ml) and IFNγ (0.1 ng/ml) stimulated ICAM1 expression eight- to ten-fold in OFs from patients with TAO (P values <0.02). Interleukin-10 (10 ng/ml) and TGFβ (1 ng/ml) had no effect. Non-specific antibody binding as assessed by the non-specific isotype control antibody, remained unchanged throughout each experimental condition.

Effect of anti-cytokine agents on cytokine responses

The anti-TNFα agents adalimumab, etanercept and infliximab inhibited TNFα-mediated ICAM1 expression in a dose-dependent manner. The IL1 receptor antagonist anakinra inhibited IL1-mediated ICAM1 expression in a dose-dependent manner. The anti-TNFα agents adalimumab, etanercept and infliximab inhibited the ICAM1 response to 0.1 ng/ml TNFα by 98% (51–144), 96% (50–143) and 90% (54–126) respectively, using peak concentrations, and 93% (50–136), 81% (53–110) and 80% (58–104) respectively, using 1/100 of peak concentrations (data expressed as mean 95% Confidence interval (CI)), P values <0.005; Fig. 1). The IL1 receptor antagonist anakinra inhibited the ICAM1 response to 0.1 ng/ml IL1 by 99% (78–119) using peak concentration and 69% (30–109) when using 1/100 of the same (P values <0.01). The effect of the anti-cytokine agents was to bring the levels of ICAM1 expression close to, or back to, baseline levels (Fig. 1).

HA measurements

OFs from seven TAO patients and four control subjects showed similar HA responses to TNFα. IL1 and their blockade (Fig. 1). In OFs from TAO patients, both TNFα (0.1 ng/ml) and IL1 (0.1 ng/ml) resulted in an increase in HA production compared with basal un-stimulated condition (for TNFα + 938 ng/ml (+44%), 95% CI 379–1498, P = 0.006 and for IL1 + 2016 ng/ml (+95%), 95% CI 1029–3003, P = 0.002; Fig. 1).

The reduction in TNFα-stimulated HA production with etanercept (2.5 μg/ml) was −1231 ng/ml (95% CI −362 to −2099, P = 0.013). The reduction in IL1-stimulated HA production with anakinra (5 μg/ml) was −1598 ng/ml (95% CI −202 to −2994, P = 0.031). This represented an inhibition of cytokine-mediated HA
production of 138% (95% CI 38–224) for etanercept and 79% (95% CI 10–149) for anakinra. The effect of the anti-cytokine agents was to bring the HA levels close to, or back to, baseline levels (Fig. 1).

**Adipogenesis**

The adipogenic media caused accumulation of lipid droplets in a proportion of OFs, visible from day 3, which increased in number and enlarged in size during the subsequent 7 days of the differentiation protocol. The ability of OFs to accumulate lipid generally decreased with increasing cell passage, so each 6- or 12-well plate contained cells from the same patient who had undergone the same number of passages, and each plate had its own control well. The mean passage number was 3 (range 1–6), and similar responses were seen in cells from patients with or without TAO.

The ORO extraction data and the data from computer analysis of digital photographs had a Pearson correlation coefficient of 0.89, \( P = 0.017 \), \( R^2 = 0.79 \) (average data from 6 plates). The ORO extraction data and the data from computer analysis of digital photographs had a Pearson correlation coefficient of 0.84, \( P = 0.038 \), \( R^2 = 0.70 \) (average data from five plates).

TNF \( \alpha \) (0.1 ng/ml) had an inhibitory effect on adipogenesis when measured by ORO extraction (\( K_{0.013} \) AU, 95% CI 0.024 to 0.002, \( P = 0.025 \), \( n = 8 \)) and visual assessment (\( K_{1.0} \), 95% CI 1.6 to 0.38, \( P = 0.007 \), \( n = 8 \)). The inhibitory effect of TNF \( \alpha \) was not significant when assessed by computer analysis of digital photographs (\( -0.02\% \), 95% CI 0.1–0.07, \( P = 0.7 \), \( n = 8 \)), but by this method, the TNF \( \alpha \) response was significantly different from the IL1 (0.1 ng/ml) response (difference of 0.56%, 95% CI 0.18–0.94, \( P = 0.01 \), \( n = 8 \)) (Figs 2–4). The inhibitory effect of TNF \( \alpha \) was attenuated by the anti-TNF \( \alpha \) agent etanercept, as measured by visual assessment in 12-well plate experiments (Fig. 3), but this did not reach statistical significance (difference = 0.56, 95% CI –0.05 to 1.12, \( P = 0.065 \)).

IL1 (0.1 ng/ml) had a stimulatory effect on adipogenesis when measured by ORO extraction (\( +0.050 \) AU, 95% CI 0.012–0.088), \( P = 0.017 \), \( n = 8 \).
visual assessment (+5.7, 95% CI 3.9–7.5, P<0.001) and computer analysis of digital photographs (+0.54%, 95% CI 0.20–0.87, P<0.007). Anakinra inhibited IL1-mediated adipogenesis measured by ORO extraction (−0.043 AU, 95% CI −0.080 to −0.005, P=0.034, n=8), visual assessment (−4.0, 95% CI −4.8 to −3.2, P<0.001) and computer analysis of digital photographs (−0.57%, 95% CI

Figure 2 Examples of adipogenesis in orbital fibroblasts from a patient with TAO. Photographs were taken with a digital camera mounted on a light microscope and lipid is stained red with oil-red-O (ORO). (A) High-power (350×) example of localized adipogenesis, the other panels are low-power (40×) photographs of orbital fibroblasts stained with ORO, with the control panel being of un-stimulated cells grown in adipogenic media for 10 days, the TNF panel being TNFα (0.1 ng/ml) in adipogenic media, and the IL1 panel being IL1 (0.1 ng/ml) in adipogenic media. Total length of scale bar represents 500 μm.

Figure 3 Effects of TNFα and IL1 and their antagonists (etanercept 2.5 μg/ml and anakinra 5.0 μg/ml) on adipogenesis, expressed as mean ± S.E.M., measured by visual assessment of microscopy appearance. A value of zero represents no adipogenesis and 30 represents maximal possible adipogenesis. Cells were grown in 12-well plates. The value labeled ‘A’ represents the un-stimulated well that contained adipogenic media but no cytokines.
representing a degree of adipogenesis was reduced across all conditions (data not shown).

**Discussion**

This study has shown that TNFα and IL1 have stimulatory effects on ICAM1 expression and GAG production, but opposite effects on adipogenesis in OFs in vitro from patients with TAO. IL1 strongly promotes adipogenesis whilst TNFα has a weak inhibitory effect.

In the absence of an accepted animal model, we used the human model of TAO, which involves culturing OFs taken from patients undergoing surgery (14, 24). This model has certain limitations. The tissue obtained at surgery is often from patients whose disease is quiescent or, if done when the disease is active, these patients are usually already on immunosuppressive treatments, such as corticosteroids. The model only comprises fibroblasts in isolation, rather than in the presence of immunologically active cells such as lymphocytes and macrophages. Despite these obvious criticisms, this model is still widely used to study OF properties.

We did not observe obvious differences, either between the behaviour of OFs from patients with or without TAO, or between OFs from patients who had or had not received corticosteroid treatment for TAO. However, our study was not designed to detect small differences between these groups. If OFs from both TAO patients and healthy controls do indeed behave in a similar way, it would suggest that it is the presence or the absence of key cytokines in the orbit that influences the occurrence of TAO, rather than the development being dependent upon inter-individual differences in OF properties.

ICAM1 is an adhesion molecule that promotes inflammatory cell recruitment when upregulated. ICAM1 has previously been shown to be present in histological sections or orbital tissue taken from patients with TAO (11, 25–27). Using immunocytochemistry IL1, TNFα and IFNγ have been shown to increase ICAM1 expression on OFs in vitro (10). In this study, we used flow cytometry to examine ICAM1 expression as it is a quantitative technique that allows more detailed study of the dose-related effects of cytokines and anti-cytokine agents. The stimulatory effects of TNFα and IL1 on ICAM1 were substantially reduced using clinically available anti-cytokine agents in concentrations achieved therapeutically. These data support the use of anti-cytokine agents to inhibit the inflammatory response in active TAO.

HA is a non-sulphated GAG that is synthesized at the plasma membrane, by a family of HA synthase enzymes. The excessive production of GAG by OFs in response to inflammatory cytokines may be one of the central mechanisms by which the autoimmune response in TAO leads to orbital tissue expansion. Orbital fibroblasts have been shown to increase their production of GAG in response to certain cytokines, such as TNFα (8), IL1 (7–9, 28), IFNγ (6, 8) and TGFβ (7). We found that both TNFα and IL1 increased HA synthesis by OFs and that these responses could be inhibited by anti-TNFα and anti-IL1 agents. These data support the use of anti-cytokine agents to inhibit GAG production and the consequent orbital tissue volume expansion in active TAO.

In addition to increased GAG production, the other main mechanism of orbital tissue volume expansion in TAO is adipogenesis. It has been shown in vitro that a
proportion of OFs have the ability to differentiate into adipocytes under certain conditions (12, 19–21). There is evidence of overexpression of mature adipocyte-related genes such as leptin, adiponectin and PPAR γ and cysteine-rich angiogenic inducer 61 in TAO compared with normal orbital tissue samples (29, 30). This supports the concept that de novo adipogenesis occurs within the orbit in active TAO.

Adipogenesis in vitro can be challenging to quantify accurately. The technique of ORO extraction is accepted and objective (23) but prone to error, particularly when the degree of adipogenesis is low due to the influence of unbound ORO. Simple visual assessment is commonly used to measure adipogenesis (12, 31). We developed a visual scoring system incorporating the number of microscopy fields containing OFs with lipid droplets, the number of cells per field containing lipid droplets and the average size of the lipid droplet. As with any other visual scoring system, it is liable to operator bias and sampling error. To reduce the influence of any operator bias we also utilized a microscopy software package system to analyse digital photographs of OFs stained with ORO. This method provided an additional objective measure of adipogenesis that correlated well with the ORO extraction and visual assessment techniques. We therefore consider that these techniques, whilst individually liable to certain errors, together are able to quantify reliably the effects of the cytokine.

Consistent with a previous study (12), we found TNFα has an inhibitory effect on adipogenesis in OFs. In our hands, the magnitude of this effect was modest, only reaching statistical significance in two out of the three methods used for quantification. This may be partly due to the relatively low levels of baseline adipogenesis that we achieved with our protocol omitting certain additives, such as carbaprostacyclin and included lower concentrations of adipogenic additives than some other studies (22). If the in vivo effect of TNFα on inhibiting adipogenesis is important in TAO, then blocking TNFα might reduce this inhibitory influence and so encourage the expansion of orbital fat. This could potentially worsen the signs and symptoms of TAO. Such an effect would be countered by the likely beneficial effects on ICAM1 expression, inflammation and GAG production, but the balance of these effects could be detrimental, particularly where adipogenesis is predominant in the disease process. It should also be considered that high-dose steroids and agents, which influence arachidonic acid metabolism, are used to promote adipogenesis in vitro, and also used as anti-inflammatory treatments for active TAO. This might mean that the pro-adipogenic properties of these agents (and possibly also anti-TNFα agents) are not clinically significant. However, the opposite view would be that the reason why the existing treatments for TAO are sometimes not as effective as desired is because of their harmful pro-adipogenic effects countering their beneficial anti-inflammatory effects.

In contrast, we found that IL1 had a strong stimulatory effect on adipogenesis. This may have important clinical implications. As IL1 promoted all three pathological aspects of the disease process studied, namely a marker of the acute inflammatory response, GAG production and adipogenesis, IL1 presents an attractive therapeutic target in TAO. Anti-IL1 agents may be the preferred anti-cytokine agent for clinical trials in active TAO both in terms of their potential to interrupt the disease processes and in terms of safety. The IL1 antagonist presently available in clinical practice (anakinra) poses many of the expected safety issues associated with an anti-cytokine agent. Nevertheless, it appears to have a more favourable safety profile compared with the anti-TNFα agents in terms of granulomatous infections, such as tuberculosis (32).

There have, as yet been no randomized, placebo-controlled trials of anti-cytokine agents in TAO, but a case series and other case reports are emerging (33, 34). These reports have used anti-TNFα agents and have generally been positive. However, the history of new treatments for TAO is sometimes of initial excitement (35), followed by an inability to repeat these positive results in larger, controlled trials (36). This may partly be due to the positive publication bias, and the fact that TAO improves spontaneously to a significant degree.

In conclusion, this study, along with others, points towards anti-TNFα agents having the potential to improve inflammation and reduce GAG production in TAO, but these beneficial effects might be counteracted by a possible worsening of any adipogenic component of the disease. In contrast, the anti-IL1 agents appear to have the potential to improve inflammation, reduce GAG production and inhibit adipogenesis, and hence could have a beneficial effect in active TAO.

Acknowledgements

This work was supported by the Health Research Board, Ireland, the Royal College of Physicians and Surgeons of Glasgow, UK and the Irish Endocrine Society.

References

5 Bahn RS. Cytokines in thyroid eye disease: potential for anticytokine therapy. Thyroid 1998 8 415–418.
The effects of TNFα and IL1 in TAO 403


9 Tan GH, Dutton CM & Bahn RS. Interleukin-1 (IL-1) receptor antagonist and soluble IL-1 receptor inhibit IL-1-induced glycosaminoglycan production in cultured human retroocular fibroblasts from patients with Graves’ ophthalmopathy. Journal of Clinical Endocrinology and Metabolism 1996 81 449–452.


18 Chang DM, Chang SY, Yeh MK & Lai JH. The pharmacokinetics of adipose conversion and triglycerides by staining intracytoplasmic lipids with Oil red O. Histochemistry 1992 97 493–497.

24 Bahn RS. The fibroblast is the target cell in the connective tissue manifestations of Graves’ disease. International Archives of Allergy and Immunology 1995 106 213–218.


Received 2 June 2006
Accepted 3 July 2006