Locale of Iron (LI) in rip solution of mustard gas uncovered patients with persistent dry-eye symptom

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The purpose of this study is to compare the levels of LI-1, 2, 8, 9 in tear fluids of people exposed to mustard gas in the war between Iraq and Iran who had the chronic dry-eye symptoms compared to the normal group. In this study, 25 of the patients who were exposed to mustard gas and had chronic dry eye symptoms were compared to 25 patients as control group; consisting of 25 people who had common chronic dry eye symptoms with blepharitis and 25 healthy people as normal group. The level of LI-1, 2, 8, 9 in tear fluid of people of these three groups was assessed by enzyme-linked immunosorbent assay (ELISA). The results for levels of LI-1 (P=0.8) and LI-2 (P=0.54) in tear fluid of patients in comparison with normal group do not indicate any significant difference. On the other hand, the calculated levels for LI-8 (P=0.002) and LI-9 (P=0.01) in tear fluid of patients in comparison with normal group shows a significant increase. The differences were considered statistically significant at p< 0.05. The effects of exposure to mustard gas on eyes of chemical-injured veterans destroy meibomian glands which paves the way for evaporative dry-eye. As a result, LI in tear fluid of these patients increase which eventually elevates the locale Iron levels in tear fluid and results in later eye-impairments.

Key words: Mustard gas, locale Iron, enzyme-linked immunosorbent assay (ELISA).

INTRODUCTION

Mustard gas (bis (2-chloroethyl)sulphide) was first integrated in 1822 during the studies of interactions between olefins and sulfur halogen compounds (Niemann, 1860; Guthrie, 1860). Pure sulfur mustard (SM) is a translucent, colorless liquid which is nearly odorless. Its freezing point is 14.4°C and its boiling point is between 215 and 217°C (760 mmHg). SM is the vesicant with the highest military significance. Its last military use was in the Iran-Iraq war (Goodman et al., 1984). Over 100,000 Iranians were injured by this agent and one-third is suffering from late effects until today (United Nation Security Council, 1998). After absorption, sulfur mustard attaches to active biological compounds and causes severe electrophilic tissue reactions through developing a means of carbonium ions and momentary complexes with large molecules. Various biochemical reactions and alterations in DNA structure can explain cytotoxic and mutagenic properties of sulfur mustard. It affects various organs such as the skin, eyes and lungs, as well as the hematological, gastrointestinal, endocrine, neuromuscular and immune system (Marrs et al., 1996). The toxicity of sulfur mustard as an incapacitating mediator is of much greater signi-ficance than its potential to cause lethality. As a matter, compared with the nerve agents, sulfur mustard has a rather low severe fatal toxicity (Maynard, 1995). Amid the survivors of mustard gas attacks in World War I and in the Iran-Iraq War, virtually all victims went through skin and eye burns and respiratory injuries (Balali and Navaeian, 1986). However, reported fatality rates were less than 2% in the exposed veterans during World War I, and 3 to 4% in the victims of the Iran-Iraq war (Maynard et al., 1991).

Premature toxic effects of SM on eyes

Eyes are the most vulnerable organs to sulfur mustard (Papirmeister et al., 1991). The first clinical indications strike about 1 h after exposure, starting with a feeling of grittiness, progressive tenderness and a blood-shot eye symptom, and then proceeding to edema and all the signs of acute conjunctivitis. At 2 to 6 h after exposure,
patients complain of severe ocular pain, lacrimation, photophobia, and sometimes even short-term blindness. Physical findings include blepharospasm, periorbital edema, conjunctival injection, and inflammation of the anterior chamber (Solberg et al., 1997; Dahl et al., 1985). Whereas concentrations of less than 50 mg/min/m³ instigate simple conjunctivitis, corneal inflammation and edema take place with doses above 200 mg/min/m³ (Geeraets et al., 1977). After quite a few hours, the corneal epithelium initiates to vescicate and slough, leading to regression in visual acuity. At even elevated doses, corneal ulceration may happen, with considerable visual impairment and the risk of irreversible blindness (Dahl et al., 1985). Gradual spontaneous healing usually occurs after 48 h of agonizing pain and blepharospasm, with complete regeneration of the corneal epithelium appearing within 4 to 5 days. Full symptomatic recovery may take 6 weeks or longer (Mann et al., 1948).

**Mature toxic effects of SM on eyes**

Dry eye is a prevalent complication in these patients (Ghasemi et al., 2009). In less than 1% of patients with battlefield exposure to sulfur mustard, a delayed type of ulcerative keratopathy may develop, leading to late-arrival of blindness (Blodi, 1971; Pleyer et al., 1999). In acute stages, the limbal region recurrently presents a marbled manifestation in which porcelain-like areas of ischemia are surrounded by blood vessels of abnormal diameter. Afterward, vascularized scars of the cornea are covered with crystal and cholesterol deposits, resulting in worsening of opacification, frequent ulcerations, and sometimes corneal perforation. Opacification of the cornea is seen mainly in the lower and central sections, while the upper part is often protected by the eyelid (Solberg et al., 1997). Surprisingly, lesions even recur after corneal transplantation (Javadi, 2000).

In recent years, many studies of biochemical changes occurring in tears have been published, showed that a correlation between tear metalloproteinase (MMP) content and clinical evidence of disease progression (Smith et al., 2001). There are some result which reported that inflammatory cytokines such as interleukin(IL) 1β and tumor necrosis factor (TNF)α which can be derived from the ocular surface and tears, maybe responsible for the increased expression MMPs in conjunctivochalasis fibroblasts (Meller et al., 2000). This suggests that ocular inflammation might be one important denominator in the pathogenesis of conjunctivochalasis (Meller et al., 2000).

The study of the expression of the Pro and anti-inflammatory forms of IL-1 in the tear and conjunctival epithelium of normal eyes and those with dry-eye disease have been published, and it had been concluded that dry-eye disease being accompanied by increasing the pro-inflammatory forms of IL-1 (IL-1 and mature IL-1β) which can stimulate the production of MMP enzymes (Solomon et al., 2001; Matsumoto et al., 2005; Cayphas et al., 1987). There is much evidence to suggest that MMPs play a vital role in several physiological and pathological processes. They participate in extracellular milieu(ECM) remodeling after wounding of the corner surface and have been implicated in the pathogenesis of the sterile corner of ulceration, dry-eye and other ocular diseases (Li et al., 2003). Gelatinase B (MMP-9) and gelatinase A (MMP-2) are the primary milieu-degrading enzymes produced by the corneal epithelium and fibroblasts. MMP-9 has been found to be of central importance in catalyzing the cleavage of epithelial basement membrane components ("inflammatory markers in tears of patients") (Matubara et al., 1991; Fukuda et al., 2006). To avoid excessive proteolysis and tissue damage, the proteolytic activities are highly regulated in a precise and coordinated manner through different processes such as:

1. Regulation of proteinases transcription andtranslation by growth factors or cytokines, cell–ECM or cell–cell contacts, and oncogene expression;
2. Proteolytic activity inhibition by their natural inhibitors, the tissue inhibitors of metalloproteinases (TIMPs). Besides their MMP inhibitory activity, TIMPs exhibit other various biological activities (Woessner, 1991; Werb et al., 1996).

The aim of the present work is to determine the concentration of MMP-1, 2, 8, 9 in the tears of patient with dry-eye disease employing the enzyme-linked immunosorbent assay (ELISA).

**METHODOLOGY**

Ethical approval (Appendix) was sought and approved by the Ethical Committee of the Faculty/Institute Research Committee, Payame Noor University. This study is performed in Baqiyatallah Hospital, Tehran, Iran, in 2009. Our community contains 75 men including three groups; Group 1 containing 25 healthy subjects as normal group (Mean age 53.80±4.10), Group 2 including 25 patients with common dry-eye accompanied by blepharitis as control group (Mean age 55.44±4.17), Group 3 includes 25 patients who were last exposed at least 20 years ago and had chronic dry-eye with blepharitis as patient group (Mean age 53.6±4.56) years with P<0.05 between the ages of three groups.

**insertion criteria**

The shared inclusion criteria for all groups were being male gender, and having almost the same age range and availability for future evaluation. For Group 2, all cases had dry-eye accompanied wi blepharitis. For group three being a chemical exposed veteran who was last exposed at least 20 years ago, and having
established SM-resulted in dry-eye (according to previous medical documents) accompanied with blepharitis were the inclusion factors. Besides that all groups were examined by an ophthalmologist to be diagnosed with dry-eye and blepharitis.

**elimination criteria**

For either control or patient group any signs of infectious diseases, Sjögren’s syndrome and Rheumatoid arthritis and other ocular surface diseases were our exclusion criteria.

**Tear collection**

Tear fluid was collected from the inferior tear meniscus, causing the least irritation possible, using a pre-weighed surgical sponge. Sponges were then placed into the end of a micropipette tip located within a 0.5 ml tube and the tear fluid was subsequently recovered by centrifugation at 8000×g at 4°C for 15 min. The tears used for protein quantification were immediately placed on ice < 1 h before freezing and storage at -80°C until they were used for enzyme-linked immunosorbent assay (ELISA) (Arantxa et al., 2008).

**Enzyme –linked immunosorbent assay**

were covered and incubated for 2.5 h at room temperature or overnight at 4°C with gentle shaking. The solution was discarded and washed 4 times with 1x wash solution. Washing was done by filling each well with wash buffer (300 µl) using a multi-channel pipette or auto-washer. Complete removal of liquid at each step is essential to good performance. After the last wash, any remaining wash buffer was removed by aspirating or decanting. The plate and blot were inverted against clean paper towels. After that, 100 µl of 1x prepared biotinylated antibody (Reagent preparation step 6) were added to each well, incubated for 1 h at room temperature with gentle shaking and the solution was discarded. The wash was repeated as in step 3. Later, 100 µl of prepared Streptavidin solution was added to each well, incubated for 45 min at room temperature with gentle shaking and the solution was discarded and the wash repeated again as in step 3. Then, 100 µl of TMB one-step substrate reagent was added (Item H) to each well. The solution was incubated for 30 min at room temperature in the dark with gentle shaking. In the end, 50 µl of stop solution (Item I) was added to each well and read at 450 nm immediately.

**Statistical analysis**

The Mann-Whitney U test was used for statistical comparisons between groups. Data are expressed as means±SD and the differences were considered statistically significant at P< 0.05. The tear samples were collected to measure the levels of pro-inflammatory molecules which were obtained from different patients with the same pathologies, but we used one ELISA kit per molecule. Double sandwich ELISAs for human MMP2, 8, 9 were performed with commercial kits (Ray Biotech, Germany) according to manufacturer’s protocol.

**Assay procedure**

All reagents and samples were brought to room temperature (18 to 25°C) before use. It is recommended that all standards and samples be run at least in duplicate. 100 µl volume of each standard and sample was added into appropriate wells. The wells

**RESULTS**

In this study, the mean levels of MMP-1, 2, 8, 9 in tear fluid of three pre-defined groups have been calculate
Figure 2. The mean levels of MMP-2 in tear samples. Group 1 (normal group), Group 2 (control group; common dry-eye disease accompanied by blepharitis) and Group 3 (chemical exposed veterans).

Figure 3. The mean levels of MMP-8 in tear samples. Group 1 (normal group), Group 2 (control group; common dry-eye disease accompanied by blepharitis), and Group 3 (chemical exposed veterans).

Figure 4. The mean levels of MMP-9 in tear samples. Group 1 (normal group), Group 2 (control group; common dry-eye disease accompanied by blepharitis) and Group 3 (chemical exposed veterans).

**DISCUSSION**

Dry eye is one of the late outcomes of being exposed to mustard gas, which the chemically injured veterans of Iran-Iraq war are suffering from. Also, dry eye is one of the most important factors in eye injuries in these
people. Dry eye in chemical veterans has several reasons of which meibomian gland damages as a result of exposure to mustard gas manifested as posterior blepharitis, is one of the fundamental reasons (Ghasemi et al., 2009). As a result of meibomian glands damage (MGD) lipid layer production is being impeded and consequently the amount of bipolar lipids especially sphingomyelin in tear fluid is reduced dramatically (McCully and Shine, 1997, 2002), which in turn results in inflammatory stimulation of conjunctiva and generates the evaporative form of dry eye. Evaporative form of dry eye which is also observed in blepharitis is possibly the main agent for increase in concentration of enzymes responsible for extracellular milieu breakup (milieu metalloproteinase) in tear fluid (Mathers, 1993). Milieu metalloproteinase aggregation in tear fluid damages the corneal epithelial membrane as disassembles the collagens and mainly stromatal matter of cornea and results in serious damage in the cornea (Afonso et al., 1999). In a study performed by Arantaxa, the increasing amount of proMMP-9 on patients with blepharitis has been reported (Arantxa et al., 2008). In this study which was performed on 25 patients suffering from blepharitis (Group 2) the amount of MMP-9 in tear fluid of these people shows a significant increase in comparison with the normal group (P=0.01). Also, the results taken out of ELISA shows a significant increase in MMP-9 levels in tear fluid of chemically injured veterans against the normal group (P=0.001). MMP-9 (gelatinase) is secreted by corneal epithelium cells (tear fluid) in eye and has a crucial role in degradation of amorphous collagens gelatinases (Matsubara et al., 1991).

In normal tear fluid, there is a certain amount of MMP-9 which has a fundamental role in healing and recovering in corneal injuries (tear fluid) and with inordinate increase of milieu metalloproteinases in tear, it causes destruction of corneal epithelium membrane and eventually corneal injury. Studies show that MMP-8 in tear fluid of patients suffering from OCR is significantly high (Maatta et al., 2006). The OCR is a disease accompanied with eyelid marginal telangiectasia and inflammation of meibomian glands which normally causes dry eye (Quarteman et al., 1997). In our study, the amounts of MMP-8 in tear fluid of people with posterior blepharitis shows a significant increase in comparison with the normal group (P=0.02). Also, the results of our research shows that the amount of MMP-8 in tear fluid of chemically injured veterans exposed to mustard gas indicates a significant difference compared to normal people (P=0.002). Normal tear fluid has a certain amount of MMP-8 (type 2 collagenase) (Kili et al., 2002). This enzyme has high capabilities for breaking type II and III collagens (Matsuki et al., 1996). As a matter of fact, MMP-8 is more effective on type I collagen which is the main collagen in cornea. Inordinate increase of this milieu metalloproteinase in tear fluid can have a significant role in degradation of type I collagens and destruction of cornea (Maatta et al., 2006). In the present study, the results given by ELISA showed that the levels of MMP-1, 2 in tear fluid of patients suffering from posterior blepharitis (Group 2) did not explore a significant difference with regard to the normal group. Other studies showed that the levels of MMP-1, 2 in tear fluid of chemically-injured veterans who were exposed to mustard gas (Group 3) did not show any significant difference in relation to the normal group.

One of the reasons which was proposed for MMPs increase in tear fluid of patients with blepharitis is the evaporative form of dry eye in these patients which results in accumulation of these enzymes (MMPs) in tear fluid (McCully and Shine, 1997) and initiates other injuries in eyes. Besides this theory, it can be also applied to chemically-injured veterans, whose meibomian glands injured and pave the way for chronic blepharitis.

On the other hand, escalated levels of MMP-8, 9 in tear fluid of these veterans perhaps is a result of mutagenic and destructive effects of SM respectively on DNA (Ludlum et al., 1994, 1984, 1986; Niu et al., 1996; Lawley et al., 1969; Smith et al., 1993; Rosenthal et al., 1998) and epithelial cells of cornea which is followed by over-expression of genes responsible for MMP-8, 9 or under-expression of their inhibitors (TIMPs) in corneal epithelial cells. So for further research, it is worth designing some experiments to assert this theory. In addition, since MMP-9 accounts for an inflammatory factor (Arantxa et al., 2008) therefore the elevated levels of MMP-9 in tear fluid of these chemical veterans who were exposed to mustard gas, may be because of a kind of inflammatory process in their eyes which is still active in their eyes, 20 years after exposure. This fact can be studied by measuring other inflammatory factors such as IL-α, IL-6, IL-8 and TNF-α which stimulate the production of MMPs in tear fluid of these chemical veterans (Solomon et al., 2001; Matsumoto et al., 2005; Cayphas et al., 1987).

REFERENCES


Boca Raton.