THE EFFECTS OF KEFIR ON THE INFLAMMATORY STATUS AND THYROID FUNCTION (EXPERIMENTAL STUDY ON WISTAR RATS AFTER EXPOSED TO CHLORPYRIFOS)

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ABSTRACT

Objective: Goiter is an enlarged thyroid gland remains a health problem in the agricultural areas. Chlorpyrifos (CPF) is a pesticide widely used by farmers. Previous studies proved that CPF exposure caused thyroid dysfunction. The objective of this study was to evaluate the effects of kefir on the inflammatory status and thyroid function in male Wistar rats after exposed to CPF using biochemical and histopathological assays.

Methods: Male rats were divided into 4 groups, i.e., CPF 5+kefir (5 mg/kg+3.6 ml/200 g, respectively), CPF 5 (5 mg/kg), corn oil (CO 1 ml/200 g), and negative control (NC: Without CPF, CO, and kefir).

Results: Kefir supplementation dose 3.6 ml/200 g once a day for 28 days in the rats after exposed to CPF dose 5 mg/kg once a day for 14 days, in CPF 5+kefir as compared to CPF 5: Significantly (p<0.05) decreased serum tumor necrosis factor-α (TNF-α) level; significantly (p<0.01) maintained serum levels of transforming growth factor-β (TGF-β) and thyroid stimulating hormone (TSH) not to decrease; not significant (p>0.05) decreased the level of interleukin-1β (IL-1β), cluster of differentiation-26 (CD26) expression and level of Tg serum; not significant (p>0.05) maintained the level of anti-thyroid peroxidase not to decrease; and not significant (p>0.05) increased the apoptosis index. This study suggests that CPF exposure causes the inflammatory process which leads to thyroid dysfunction.

Conclusion: Kefir supplementation significantly decreased the level of TNF-α and maintained the levels of TGF-β and TSH not to decrease, possible to reduce the inflammatory and thyroid dysfunction processes caused by exposure to CPF in experimental animals.

Keywords: Kefir, Chlorpyrifos, Inflammation, Thyroid function.

INTRODUCTION

High prevalence of goiter, especially on primary school (PS) children with normal urinary iodine excretion (UIE) remains a health problem in agricultural areas. In working area of Kluwut Primary Health Care, Bulakamba Subdistrict, total goiter rate (TGR) in PS children had increased from 2012 to 2014 (32.17%, 48.97%, and 50.46%, respectively). Data from Brebes District Health Office in 2010 showed UIE in PS children in Kluwut was 286-293 μg/l [1]. According to WHO, UIE ≥100 μg/l was categorized as adequate [2]. The TGR in Kluwut was very high compared to a national survey in 2003, which showed TGR in PS children was 11.2% and median UIE 229 μg/l [3]. The WHO determines an area with TGR ≥30.0% is categorized as high endemic. The WHO had increased from 2012 to 2014 (32.17%, 48.97%, and 50.46%, respectively). Data from Brebes District Health Office in 2010 showed UIE in PS children in Kluwut was 286-293 μg/l [1]. According to WHO, UIE ≥100 μg/l was categorized as adequate [2]. The TGR in Kluwut was very high compared to a national survey in 2003, which showed TGR in PS children was 11.2% and median UIE 229 μg/l [3]. The WHO determines an area with TGR ≥30.0% is categorized as high endemic. The WHO had increased from 2012 to 2014 (32.17%, 48.97%, and 50.46%, respectively). Data from Brebes District Health Office in 2010 showed UIE in PS children in Kluwut was 286-293 μg/l [1]. According to WHO, UIE ≥100 μg/l was categorized as adequate [2]. The TGR in Kluwut was very high compared to a national survey in 2003, which showed TGR in PS children was 11.2% and median UIE 229 μg/l [3]. The WHO determines an area with TGR ≥30.0% is categorized as high endemic. The WHO had increased from 2012 to 2014 (32.17%, 48.97%, and 50.46%, respectively). Data from Brebes District Health Office in 2010 showed UIE in PS children in Kluwut was 286-293 μg/l [1]. According to WHO, UIE ≥100 μg/l was categorized as adequate [2]. The TGR in Kluwut was very high compared to a national survey in 2003, which showed TGR in PS children was 11.2% and median UIE 229 μg/l [3]. The WHO determines an area with TGR ≥30.0% is categorized as high endemic.

Exposure to CPF has been reported differently by several researchers that CPF exposure has caused thyroid follicular cells proliferation, necrosis, or apoptosis [4-6]. Previous studies proved that CPF exposure could cause an increasing or a decreasing thyroid stimulating hormone (TSH) and thyroxine hormone (T₄) levels [6-10]. We assumed that CPF affected thyroid dysfunction, but it was not clear whether the CPF affected hypothyroidism or hyperthyroidism and have not reported a possible the inflammatory process in the pathogenesis of thyroid dysfunction caused by CPF exposure.

The cells apoptosis result of CPF exposure may affect to other cells. Although apoptosis appears to play a role in the pathogenesis of both Hashimoto's thyroiditis (HT) and Graves' disease (GD), the mechanisms that mediate, these processes appear different. The induction of apoptosis in HT results in the destruction of thyrocytes, while apoptosis in the GD leads to damage of thyroid-infiltrating lymphocytes. The differences in the apoptotic mechanisms produce two very different forms of thyroid autoimmune responses, eventually developing into HT and GD, respectively [11]. Thyroid follicular cells apoptosis can be sensitized by pro-inflammatory cytokines interferon-γ (IFN-γ) and interleukin-1β (IL-1β) or tumor necrosis factor-α (TNF-α) and IL-1β through the mediation of Fas ligand or TNF-related apoptosis-inducing ligand [12-14]. Previous studies proved that the levels of TNF-α and IL-1β increased also correlated with GD [15-18].

Previous studies reported that exposure to CPF increased the levels of TNF-α and IL-1β, increased the production of IL-1β after induced by lipopolysaccharide and the expression of a cluster of differentiation-26 (CD26) as well as the levels of IL-10 and transforming growth factor-β (TGF-β) [19-25]. Based on this description, it can be concluded that CPF exposure induces pro-inflammatory cytokines TNF-α, IFN-γ, and IL-1β. Apoptosis in thyroid follicular cells or thyroid-infiltrating lymphocytes on the pathogenesis of thyroid dysfunction are possible through sensitization of these pro-inflammatory cytokines caused by CPF exposure.

Kefir has a role as anti-inflammatory and anti-apoptosis. Lactic acid bacteria in kefir modulate the immune system to produce anti-inflammatory cytokines IL-10 and TGF-β [26-28]. IL-10 acts as an immunosuppressive through increases the life expectancy of the cells by increasing the production of anti-apoptosis B-cell lymphoma-2 (Bcl-2) [29]. The effects of anti-inflammatory IL-10 are thought to be caused by a reduced production of the pro-inflammatory cytokines IL-12, IFN-γ and TNF-α while TGF-β regulates the adaptive immune system directly inhibit T-cell proliferation by reducing the production of IL-2 and repressing level of IL-1β induced proteinase-activated receptor-2 expression. IL-10 and TGF-β also play a role in therapy in autoimmune disease models [27,29-31].
Kefir has a role as a regulator as anti-apoptosis and pro-apoptosis. The role of anti-apoptosis kefir is to reduce the levels of pro-apoptotic protein Bcl-2-associated-x (Bax), Bcl-2-associated agonist of cell death (bad), sitokom c, caspase-3, and decrease the level of the pro-inflammatory cytokine TNF-α [32]. Lactobacillus rhamnosus GG on kefir activates the anti-apoptotic Akt/protein kinase B. This model protobiotic also inhibits activation of the pro-apoptotic p38/mitogen-activated protein kinase (MAPK) by TNF-α, IL-1α, and INF-γ [33]. The role of pro-apoptosis kefir is to upregulate the ratio of protein bax/Bcl-2 and increase the expression of p53 independent p21 expression. The apoptosis increases with increasing kefir concentration. Western blot analysis demonstrates that kefir induces overexpression of bax, while repressing Bcl-2 [34,35]. Previous studies have not reported the role of kefir as anti-inflammation in the case of thyroid dysfunction caused by CPF exposure. This study aimed to investigate the efficacy of anti-inflammatory properties of kefir on Wistar rats after exposed to CPF causing thyroid dysfunction.  

METHODS  

Animals  
This study was carried out by using male Wistar rats (250-400 g). Rats were obtained from the Integrated Research and Development Institute of Unit IV (LPPT) Gadjah Mada University in Yogyakarta. The animals were maintained under standard laboratory conditions (temperature 27-32°C) with dark and light cycle (12/12 hrs) and allowed free access to standard pellet diet (The National Agency of Drug and Food Control, Indonesia) and water ad libitum. The rats were acclimatized to laboratory condition for 1 week before the commencement of experiment. All procedures described conducted in accordance with Guideline for Care and Use of Animals Laboratory of LPPT. This study was approved by the Research Ethics Committee for Health Research No. 62/EC/FK-RSDK/2015, Faculty of Medicine, Diponegoro University and Dr. Kariadi General Hospital, Semarang, Indonesia.

CPF, corn oil (CO) and kefir  
CPF (Dursban® 200 EC) was obtained from Store for Trading Pesticides and Agricultural Apparatus in Brebes, Indonesia. Dursban® 200 EC examined using gas chromatography method was proven that this insecticide containing CPF 218.5 g/l. Before Dursban was given to experimental animals orally by gavage, it was dissolved in CO for a final concentration of 0.5 mg/ml. CO was obtained from local supermarkets in Brebes, Indonesia in the form of refined CO with plastic packaging. Kefir was made from the 24 hrs fermented cow milk by kefir grains commercial inoculum that obtained from the House of Kefir Ungaran, Semarang, Indonesia.

Experimental study  
Randomized control group pre- and post-test design performed on 36 Wistar rats were divided into 4 groups, i.e., (1) CPF 5+kefir and, (2) CPF 5, rats were coadministered CPF 5 mg/kg (1/5 LD50), (3) CO, rats, were coadministration CO 1 ml/200 g. CPF (dissolved in CO) and CO were given orally by gavage once a day for 14 days. Kefir was given from the 15th to 42nd days (28 days) dose 3.6 ml/200 g orally by gavage once a day. Pre- and post-test data were measured on 14th and 42nd days, including levels of TNF-α, IL-1β, and TGF-β; expression of CD26; levels of TSH, T3, and antithyroid peroxidase antibody (anti-TPO). One rat of each group was sacrificed for histopathology examination for hematoxylin and eosin (H and E) staining and immunohistochemical with caspase-3 staining (thyroid follicle cells apoptosis index). Four negative control (NC), the data in this study to determine the value that was considered normal were also taken as above in rats without treatment of CPF, CO and kefir as an NC. The data were measured only at the time of the post-test.

Evaluation of the inflammatory status and thyroid function  
Cytokines (TNF-α, IL-1β, and TGF-β), hormones (TSH and T3) and anti-TPO concentrations were assayed using double-antibody sandwich enzyme-linked immunosorbent one-step process assay kit (Qayee-Bio, Shanghai China). CD26 kit using rat CD26 (OX61); Sc-53039 Santa Cruz Biotechnology Inc. USA and CD26 expression was analyzed using flow cytometer.

Histopathological study  
Skin of the neck was incised, and trachea was removed. Thyroid gland on the posterior aspect of trachea was removed. They were washed in saline, dried by a filter paper and then they were fixed in 10% neutral buffered formal and processed for paraffin blocks. 5 µm paraffin sections were cut and stained with H and E stain for routine histopathological study.

Immunohistochemical stains for thyroid sections  
The rabbit polyclonal anti-active/cleaved caspase-3 antibody (Novus Biologicals, Littleton, Colorado, USA) was used. To ensure antibody specificity, NC samples were processed under the same conditions but without using the primary antibody. Brown color in the cytoplasm was considered a positive reaction. The percent area of positive cells was measured using image analysis in five randomly selected, separate, ×400 magnified fields from each slide.

Statistical analysis  
The data were presented as the mean±standard deviation. The differences pre- and post-test were analyzed using Paired t-test or Wilcoxon. The differences among the three groups with NC were analyzed using independent t-test or Mann-Whitney. The statistical significance of differences between the groups was assessed with a one-way ANOVA or Kruskal-Wallis test, followed by Duncan post-hoc test analysis.

RESULTS  

The effect of kefir on the inflammatory status  

The effect of kefir on serum TNF-α level  
The effect of kefir on serum TNF-α level of rats after to exposed CPF is presented in Fig. 1A. Pre-test data showed TNF-α level in CPF 5+kefir and CPF 5 higher than NC rated significant (p<0.05), but in CO lower than NC not significant (p>0.05). TNF-α level after 28 days (post-test) significantly decreased (p<0.05) only in CPF 5+kefir and CO, but on CPF 5 decreased not significant (p>0.05). Test of delta (Δ) TNF-α level (Table 1) with ANOVA in three groups rated significant (p<0.05). post-hoc analysis with Duncan test showed that decreasing level of TNF-α in CPF 5+kefir more than CPF 5. In CPF 5 decreased fewest.

The effect of kefir on serum IL-1β level  
Pre-test data (Fig. 1B) showed IL-1β level in CPF 5+kefir, CPF 5, and CO higher than NC, but not significant (p>0.05). Post-tests data showed IL-1β level in CPF 5+kefir decreased not significant (p>0.05), but on CPF 5 and CO increased not significant (p>0.05). Test of delta (Δ) IL-1β level (Table 1) with ANOVA in three groups rated not significant (p>0.05). post-hoc analysis with Duncan test showed that decreasing level of IL-1β most differs occurred in the group CPF 5+kefir.

The effect of kefir on serum TGF-β level  
Pre-test data (Fig. 1C) showed TGF-β level in CPF 5+kefir, CPF 5, and CO higher than NC, but not significant (p>0.05). Post-test data showed TGF-β level in CPF 5 decreased significant (p<0.05) in CPF 5+kefir, decreased not significant (p>0.05) whereas in CO group TGF-β level increased not significant (p>0.05). Test of delta (Δ) TGF-β level (Table 1) with ANOVA in three groups rated not significant (p>0.05). post-hoc analysis with Duncan test showed that decreasing TGF-β level in the most different in CPF 5, whereas in CPF 5+kefir decreased fewest.

The effect of kefir on serum CD26 expression  
Pre-test data (Fig. 1D) showed CD26 expression in CPF 5+kefir significantly (p<0.001), CPF 5 (p<0.001) and CO (p<0.01) higher than NC. Post-test data showed that expression of CD26 in CPF 5+kefir significantly (p<0.001), CPF 5 (p<0.01) and CO (p<0.01) decreased. Δ
CD26 expression in CPF 5+kefir decreased more than CPF 5. Test of Δ CD26 expression (Table 1) with ANOVA in three groups rated not significant (p>0.05). Post-hoc analysis with Duncan test showed that decreasing CD26 expression most differs occurred in CO, but in CPF 5+kefir decreased more than CP 5. In CPF 5 decreased fewest.

The effect of kefir on the thyroid function

The effect of kefir on serum TSH level

The effect of kefir on serum TSH level of rats after exposed to CPF is presented in Fig. 2A. Pre-test data showed TSH level in CPF+kefir, CPF 5 and CO lower than NC, but not significant (p>0.05). Post-test data showed TSH level in CPF+kefir, CPF 5 and CO decreased not significant (p>0.05). Test of Δ TSH level (Table 1) with Kruskal-Wallis in three groups rated significant (p<0.01). Δ TSH level most decreased in CPF 5+kefir then CPF 5 and CO. In CO decreased fewest.

The effect of kefir on serum T4 level

The effect of kefir on serum T4 level of rats after exposed to CPF is presented in Fig. 2B. Pre-test data showed T4 level in CPF+kefir, CPF 5 and CO not significantly higher than NC (p>0.05). Post-test data showed T4 level in CPF+kefir, CPF 5 and CO decreased not significant (p>0.05). Post-test T4 level in CPF 5+kefir is almost the same as NC, while in the other two groups are still higher than NC (p>0.05). Test of Δ T4 level (Table 1) with ANOVA in three groups was not significant (p>0.05). Δ T4 level most decreased in CPF 5+kefir then CPF 5 and CO. In CO decreased fewest.

The effect of kefir on serum anti-TPO level

The effect of kefir on serum anti-TPO level of rats after exposed to CPF is presented in Fig. 2C. Pre-test data showed the anti-TPO level in CPF+kefir and CPF 5 higher than NC, but in CO lower than NC (p>0.05). Post-test data showed the anti-TPO level in CPF+kefir, CPF 5 and CO decreased not significant (p>0.05). Test of Δ (Table 1) the anti-TPO level with ANOVA in three groups rated not significant (p>0.05). Post-hoc

### Table 1: Summary data of changes (Δ) in the value of various variables among groups of experimental animals

| Variables       | Experimental animal groups | p    |
|-----------------|----------------------------|------|
|                 | CPF5+Kefir                 | CPF5 | CO   |
| TNF-α (ng/ml)   | −14.93±10.16               | −11.46±12.46 | −39.93±18.15 | 0.011 g* |
| IL-1β (pg/ml)   | −7.44±7.73                 | 0.65±5.14 | 1.08±10.00 | 0.375 g |
| TGF-β (ng/ml)   | −1.17±4.37                 | −5.28±3.96 | 5.89±6.03 | 0.008 g** |
| CD26 (%)        | −44.07±8.71                | −42.32±2.22 | −47.43±13.77 | 0.008 g** |
| TSH (ng/ml)     | −3.26±12.92                | −35.37±18.10 | −39.02±25.54 | 0.009 h** |
| T4 (ng/ml)      | −5.44±8.83                 | −3.93±15.09 | −1.33±12.56 | 0.891 g |
| Anti-TPO (ng/ml)| −1.91±12.33                | −3.03±6.13 | −3.34±8.61 | 0.912 g |
| Apoptosis index (%) | 5.00±1.41                 | 0.50±2.12 | −4.50±0.71 | 0.102 h |

Data were expressed as mean of Δ±SD for seven rats in each group. g: One-way ANOVA test; h: Kruskal-Wallis test; *p<0.05, **p<0.01, ***p<0.001, significantly different to each groups. CPF: Chlorpyrifos, CO: Corn oil, TNF-α: Tumor necrosis factor-α, IL-1β: Interleukin-1β, TGF-β: Tumor growth factor-β, TSH: Thyroid stimulating hormone, TPO: Thyroid peroxidase, Δ: Delta
analysis with Duncan test showed decreasing the anti-TPO level in the most different in CPF 5 and CO, whereas in CPF 5-kefir decreased fewest. In CPF 5+kefir decreased fewest.

**The effect of kefir on serum apoptosis index**

The effect of kefir on the apoptosis index of rats after exposed to CPF is presented in Fig. 2D. Post-test data showed apoptosis index in CPF 5-kefir, and CPF 5 increased not significant (p<0.05), whereas in CO decreased not significant (p>0.05). The apoptosis index in CPF 5-kefir increased more than CPF 5. Test of Δ (Table 1) the apoptosis index in three groups with Kruskal-Wallis in this study rated not significant (p>0.05). Test of Δ (Table 1) the apoptosis index with Kruskal-Wallis in three groups were not significant (p>0.05). Δ the apoptosis index most increased in CPF 5+kefir, but in CO decreased.

**DISCUSSION**

The result of this study showed similar with the previous study that exposure to CPF increased the level of TNF-α [19,20]. The previous study revealed significant increase of TNF-α secretion among CPF-exposed workers. Enhancement of TNF-α release was explained by the fact that insecticides modulate immune response via different mechanisms: Thl-like immune response was enhanced with the release of cytokines (IL-2 and TNF-α) affecting B-cell maturation and immunoglobulin production. The IL-2 and TNF-α increase may result from a mechanism to compensate for the decrease in TNF-γ after Insecticide exposure [19]. Protein kinases like the Akt family or extracellular signal-regulated kinase (ERK), which are essential for cell survival and proliferation, were inhibited by CPF [22]. Kefir supplementation dose 3.6 ml/200 g significantly reduced level of TNF-α in Wistar rats after exposed to CPF 5 mg/kg. Kefir modulated the immune system to produce anti-inflammatory cytokines IL-10 [26,28]. IL-10 promoted the development of a Type 2 cytokine pattern by inhibiting the IFN-γ production of T lymphocytes directly inhibited the proliferation of CD4+ T-cells and production of cytokines such as IL-2, IFN-γ, IL-4, IL-5, and TNF-α [29]. Kefir also contains Saccharomyces cerevisiae [36]. In vitro studies, S. cerevisiae var. boulardii role in blocking the activation of nuclear factor kappa B (NF-kB) and MAPK decreases the expression of inflammation-associated cytokines, such as IL-8, TNF-α, and IFN-γ. Anti-inflammatory effect of S. cerevisiae var. boulardii exerts anti-inflammatory effects after stimulation with Clostridium difficile-toxin A due to decrease in secretion of IL-8 in human colonocytes and activation of ERK 1/2 in both human colonocytes and murine ileal loops also decreased level of IL-8 in human colon and activated signal ERK 1/2 in human colon and ileum murine [36,37]. IL-10, expressed by macrophages and dendritic cells, is dependent on the activation of ERK 1/2 [22].

This study similar with previous study that exposure to CPF increased the level of IL-1β [22,23]. The previous study revealed that several stress stimuli (chemical stress, osmotic shock, heat shock) induced 5-lipooxygenase product formation in freshly isolated polymorphonuclear leucocytes, in parallel with activation of p38 MAPK [38]. Leukotriene B4, a major product of 5-lipoxygenase, has been shown to augment the IL-1β release [39]. Exposure to CPF increases the level of IL-1β in the rats serum, possible through an increasing the enzyme 5-lipooksigenase who played a major role in producing leukotriene B4. Kefir supplementation dose 3.6 ml/200 g reduced level of IL-1β after exposed to CPF 5 mg/kg, but not significant (p>0.05). Kefir modulated the immune system to produce anti-inflammatory cytokine TGF-β [26,28]. TGF-β acts as an inhibitor of the actions of IL-2 and IL-1β [29,30,40]. Kefir also contains Lactobacillus helventicus [36]. The oral administration of L. helventicus HY’7801 in Candida albicans-infected mice induced a reduction of NF-kB activation in the vaginal tissue, decreased the expression of IL-1β, TNF-α, IL-6, cyclooxygenase-2, and inducible nitric oxide synthase, and increased the expression of IL-10. The role of L. helventicus Hc-10 in reducing the production of IL-1β by inhibiting the activity of leukotriene B4 has been proved increasing the level of IL-1β [39,41].
This study is consistent with previous study that exposure to CPF reduced the level of TGF-β [25]. Kefir supplementation 3.6 ml/200 g for 28 days significantly maintained the TGF-β level not to decrease (relative increasing) after exposed to CPF 5 mg/kg. TGF-β is produced by Th3 cells mainly in the gastrointestinal mucosa; therefore, Th3 cells are very important in maintaining the tolerance of antigens orally [42]. The ability of kefir as probiotics maintained the level of TGF-β not to decrease after exposure to CPF because kefir has been shown to improve homeostasis gastrointestinal tract. The interaction between probiotic strains and enterocytes is important for the controlled production of cytokines and chemokines secreted by epithelial cells. Indeed, it has been shown that some probiotic organisms can modulate the in vitro expression of pro and anti-inflammatory molecules in a strain-dependent manner. For instance, Lactobacillus sakei induces the expression of IL-1β, IL-8, and TNF-α, whereas Lactobacillus johnsonii stimulates the production of TGF-β in caco-2 cells. This process appears to require cross-talk between the epithelial cells and the underlying leukocytes [27,43].

The result of this study is consistent with previous study that exposure to CPF in this study increased the CD26 expression [24]. Exposure to CPF increases level of IL-2 [19]. The previous study reported that both IL-2 and IL-12 up-regulated the expression of adenosine deaminase and CD26 [44]. High CD26 cell surface expression is correlated with the production of Th1-type cytokines such as IFN-γ [45], it is correlated with the production of TNF-α and IL-β. Kefir supplementation 3.6 ml/200 g decreased expression of CD26 after exposure to CPF dose 5 mg/kg but not significant. The role of kefir is to inhibit the activity of CD26 through β-lactoglobulin which is produced by Lactococcus lactis and through the ability to decrease the level of IL-2 [46,47]. In this study, the ability of kefir in decreasing expression of CD26 is correlated with the ability to decrease the levels of TNF-α and IL-β, which increased after exposure to CPF and maintain TGF-β level not to decrease.

The previous study proved that exposure to CPF decreased level of TSH [9,10]. In this study, post-test data showed that the TSH level decreased in conjunction with increasing the TNF-α and IL-β levels possible were caused by the inflammatory processes in central nervous system. It is suspect that exposure to CPF may cause inflammation on the pituitary or hypothalamus gland. Previous study showed that the monocrotophos (organophosphate) pesticide disturbed the thyroid hormones (THs) homeostasis and interfered with the transport and conversion of THs, synthesis and secretion of pituitary TSH, and regulation of hypobalamic thyrotropin-releasing hormone (TRH) in female goldfish [48]. Obstacles the activity of acetylcholinesterase (AChE) enzymes caused by organophosphate exposure will increase the acetylcholine level. After cholinergic activation, leading to somatostatin release TRH secretion will be inhibited. There is evidence that acetylcholine is involved in regulating pituitary functions. Several lines of evidence support a role for cholinergic regulation of TSH secretion through by somatostatin. Dopamine stimulates somatostatin release from the median eminence and increases its portal blood concentration, and this increase suppresses the serum TSH level [49]. Organophosphate exposure, even if there are no obstacles the activity of AChE enzymes, still can induce inflammation through increased production of cytokines such as TNF-α, IL-1β, and IL-6 in the cortex, hippocampus, and thalamus of rats [50]. The toxic effect of CPF on the central nervous system which reduced the TSH level could be through the inflammatory processes. Kefir supplementation significantly maintained (relative increasing) the TSH level not to decrease after exposure to CPF, possible caused the effects of kefir as an anti-inflammation to reduce the activity of cytokines such as TNF-α, IL-1β in the brain.

This study is similar with previous study that exposure to CPF tends to increase the T₄ level [10]. Kefir supplementation dose 3.6 ml/200 g for 28 days in Wistar rats tends to decreased level of T₄ after exposed to CPF 5 mg/kg. Based on this study, exposure to CPF caused decreasing the level of TSH and tends to increase the level of T₄ so thyroid dysfunction in this study was likely to cause hyperthyroidism. The previous study has proved that increasing the levels of TNF-α and IL-1β correlated with GD. In addition, IL-1β induces production of hyaluronan by primary thyroid epithelial cells and thyroid fibroblasts, a process that may contribute to the development of goiter in GD [15-18].

High levels of IL-1β and low level of TGF-β can be correlated with high levels of autoantibodies. In the development of GD, infiltration of the thyroid by activated immune cells results in local release of IL-1β. It has been observed that IL-1β induces the production of IL-6, IL-8, intercellular adhesion molecule-1, and other inflammatory mediators. IL-1β also enhances T cell-dependent antibody production by augmenting CD40 ligand and OX40 expression on T-cells. IL-1β was shown to promote differentiation of T-helper 1 (Th1) cell, the proportion of which was reported to be higher in intractable GD than that of GD in remission [15]. In fact, patients with autoimmune diseases, such as systemic lupus erythematosus (SLE), have reduced TGF-β production in their peripheral blood cell cultures. Hence, reduced TGF-β production by immune cells might predispose to autoactive T-cell activation and autoantibody production in autoimmune diseases [51]. CD26 is a multifunctional ectoenzyme involved in T-cell activation that has been implicated in autoimmune pathophysiology. CD26 may contribute to the orchestration of the immune response by Th17 cells in human inflammatory diseases. IL-17-producing CD4+ T-cells (Th17 cells) are important mediators of autoimmune disease [52].

In vitro study, CPF exposure in JEG-3 choriocarcinoma cell line modulated the mRNA levels of pro-inflammatory IL-6, IL-17 and the anti-inflammatory IL-13 cytokines [53]. IL-17 promotes the survival
and expansion of B-cells and the differentiation of B-cells into antibody-producing plasma cells. This may lead to autoimmunity. IL-17 also induces the production of other pro-inflammatory cytokines, such as TNF-α and IL-1β [54]. Antibodies against thyroid antigens are present in GD and HT diseases, but their specific epitopes are different, resulting in different functional antibodies. Autoimmune thyroid disease development occurs due to loss of immune tolerance to autoantigens and reactivity of the thyroid, which leads to infiltration of the gland by T-cells and B-cells, which in turn produce antibodies specific for clinical manifestations of hyperthyroidism and hypothyroidism, GD and HT, respectively. IL-17 is secreted by Th17 cells; IL-17 play an important role in chronic inflammatory diseases such as asthma and SLE. The percentage of Th17 lymphocytes in GD patients not treated with anti-thyroid drugs higher than GD remission patients [55].

In this study, the levels of TNF-α, IL-1β, and CD26 expression increased after exposure to CPF and the level of TGF-β decreased in conjunction with increasing the inflammatory process on the thyroid gland (Fig. 3A), but not accompanied with increasing the level of anti-TPO (likely to fall). Kefir supplementation in this study tends to maintain anti-TPO level not to decrease after exposure to CPF, but not significant. Based on this study, exposure to CPF caused increasing the levels of TNF-α, IL-1β, and CD26 expression may be associated with increasing the other autoantibodies, especially TSH receptor antibody. This study is consistent with the previous study that the incidence of thyroid dysfunction after exposed to CPF significantly decreased the TSH level and was about to tend to increase the T₄ level (hyperthyroidism) [10].

The newest case of organophosphorus (CPF) intoxication was reported and was about to tend to increase the T₄ level) [56]. The role of kefir decreasing the level of T₄ is very likely related to the role of kefir as immunoregulatory by balancing the pro-apoptosis in thyroid follicular cells and anti-apoptosis in the process of thyroid-infiltrating lymphocytes, as well as the role of anti-inflammatory to reduce the levels of TNF-α and IL-1β and maintain the level of TGF-β. Kefir as immunoregulator can act as an anti-apoptosis and pro-apoptosis agent. In this study, the role of kefir enhances the apoptosis index (pro-apoptosis) on CPF5-kefir group (Fig. 3f) possible to decreased or normalized the level of T₄ (almost equal with NC) after exposure to CPF. Lactic acid bacteria in kefir modulate the immune system to produce anti-inflammatory cytokines IL-10 and TGF-β [26-28]. IL-10 and TGF-β, exhibit suppressive activities on macrophage functions and antagonize the effect of Th1-secreted cytokines. All of this may reflect the complexity of the feedback regulatory mechanisms that occur through the cytokine network in macrophages. IL-10 also exerted a suppressive effect on IL-17-induced TNF-α release, while the inhibitory effects of IL-4, IL-13, and TGF-β2 on TNF-α secretion were partial [57].

The data suggest a pivotal role for kefir in combating inflammatory and autoimmune responses are thought to be caused by CPF exposure.

**CONCLUSION**

The present study suggests that CPF exposure causes the inflammation through increasing the levels of TNF-α, and decreasing the level of TGF-β. This inflammatory processes may interfere the pituitary-thyroid axis which causes thyroid dysfunction by decreasing the serum T₄ level and tend to increasing the serum T₃ level (hyperthyroidism). Based on this study, the role of kefir as an anti-inflammatory is through the ability to decrease the serum level of TNF-α, maintain the levels of TGF-β and TSH not to decrease and tend to decrease the serum T₄ level possible to cure thyroid dysfunction (hyperthyroidism) in the rats caused by exposure to CPF.

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