Selenium supplementation protects trophoblast cells from oxidative stress

Author
Watson, Marlee, Van Leer, Lyndal, Vanderlelie, Jessica, Perkins, Anthony

Published
2012

Journal Title
Placenta

DOI
https://doi.org/10.1016/j.placenta.2012.09.014

Copyright Statement
Copyright 2012 Elsevier Ltd. This is the author-manuscript version of this paper. Reproduced in accordance with the copyright policy of the publisher. Please refer to the journal's website for access to the definitive, published version.

Downloaded from
http://hdl.handle.net/10072/51829
Selenium Supplementation Protects Trophoblast Cells from Mitochondrial Oxidative Stress.

Alisha Khera, Jessica J. Vanderlelie, Anthony V. Perkins
School of Medical Science, Griffith Health Institute,
Griffith University, Gold Coast Campus
Southport, Queensland, Australia.

Corresponding author:
Professor Tony Perkins
School of Medical Science,
Griffith University Gold Coast Campus
Parklands Drive, Southport,
QLD, 9726, Australia

E-mail: a.perkins@griffith.edu.au
Phone: + 61 (0) 7 555 28818
Fax: + 61 (0) 7 555 29081

Keywords: Trophoblast, Oxidative Stress, Selenium, Reactive Oxygen Species, Antioxidants
ABSTRACT

INTRODUCTION: Oxidative stress plays an important role in the pathogenesis of preeclampsia, a placental disorder affecting approximately 7% of pregnancies. Trophoblast cells are susceptible to oxidative stress which causes increased cell death and placental turnover. In this study, inhibitors of the mitochondrial respiratory chain were utilised to induce oxidative stress and the effect that selenium supplementation had on trophoblast viability was investigated.

METHODS: Trophoblast cells (BeWo, JEG-3 and Swan-71) were treated with Na Selenite (100nM) or Selenomethionine (500nM) to increase the biological activity of antioxidants Glutathione Peroxidase and Thioredoxin Reductase. The cells were then oxidatively stressed with the addition of increasing doses of Antimycin C and Rotenone and the Resazurin end point assay was used to assess cellular activity.

RESULTS: There was a significant dose dependent decrease in the cellular activity in BeWo, JEG-3 and Swan-71 when treated for 4 hours with increasing concentrations of Antimycin (40-320µM) and Rotenone (100-800nM). Prior Incubation with Na Selenite and Selenomethionine was able to protect trophoblast cells from oxidative stress at Rotenone concentrations of 200nM and 400nM (P<0.001) and Antimycin concentrations of 80-240 µM (P<0.001).

DISCUSSION: These data suggest that selenoproteins such as Glutathione Peroxidase and Thioredoxin Reductase have an important role in protecting trophoblast mitochondria from oxidative stress.

CONCLUSIONS: This study emphasises the importance of maintaining an adequate selenium supply during pregnancy and especially in pregnancies complicated by conditions such as preeclampsia.
INTRODUCTION

Oxidative stress describes a condition where the presence of reactive oxygen and nitrogen species (RONS) overwhelms cellular antioxidant defences. The initial response to oxidative stress is to up regulate defensive mechanisms such as heat shock proteins and anti-oxidant enzymes. Cells enter a state of suspended animation whilst the stress is counteracted or disperses at which point the cells return to homeostasis. If the oxidative insult persists, oxidised macromolecules such as lipids, proteins and nucleic acids are released. Persistent or excessive oxidative stress will lead to autophagy, apoptosis and necrosis with the associated release of cellular debris [extensively reviewed 1]. Placental oxidative stress is central in the pathogenesis of many disorders of pregnancy including preeclampsia and preterm labour and numerous studies have shown that there is increased shedding of oxidised macromolecules as well as cellular debris from placentae affected by these disorders [2,3].

Anti-oxidant enzymes are central to protecting cells from oxidative stress. These include enzymes involved in the Glutathione and Thioredoxin reducing systems such as Glutathione Peroxidase (GPx) and Thioredoxin Reductase (ThxRed). These two proteins are selenoenzymes, whereby activity in all cells and tissues is dependent upon an adequate supply of the essential micronutrient selenium. This provides an elegant mechanism for up-regulation, as selenium supplementation is able to promote the activity of these proteins in many cell types, in animal studies and in humans [4,5,6].

Previous work from this laboratory proposed that the up-regulation of GPx and ThxRed using selenium supplementation would protect cardiac tissues from the damaging effects of the oxidative stress during clinical procedures including coronary artery bypass grafts. This proved to be the case and in clinical studies the beneficial effects of selenium supplementation, in association with other essential micronutrients, resulted in significantly quicker patient recoveries [7]. The same approach is an attractive proposition when considering placental oxidative stress in complications of pregnancy such as preeclampsia. Could maximising placental expression of these proteins using selenium supplementation protect trophoblast from oxidative stress?
In a recently published paper we demonstrated that up-regulation of GPx and ThxRed using both inorganic sodium selenite and organic selenomethionine was able to protect trophoblast from exogenously added reactive oxygen species such as peroxides [8]. In the present study we have asked the same question but generated the oxidative stress endogenously by selectively blocking the electron transport chain and generating mitochondrial oxidative stress.

**METHODS AND MATERIALS.**

**Cell Culture and Reagents:**
The human choriocarcinoma cell lines BeWo and JEG-3 cells were obtained from American Tissue Culture Collection (ATCC). The non-choriocarcinomic trophoblastic cell line Swan-71 was kindly provided by Professor Gil Mor, Yale University. Cells were grown and maintained at 37°C with 5% CO₂ in Dulbecco’s Modified Eagle Medium (Invitrogen, Australia) containing 10% fetal bovine serum and 500 U/mL penicillin–streptomycin (Invitrogen, Australia). Antimycin and Rotenone were obtained from Sigma-Aldrich Ltd, Australia. All other reagents, unless stated otherwise, were obtained from Sigma- Aldrich Ltd, Australia. Trophoblast cells were collected from 75cc flasks by trypsinization and were seeded in 96 well sterile tissue culture plates at 10,000 cells per well for treatment and analysis.

**Cell Extraction:**
To extract protein for biochemical analysis, the confluent monolayer from 75cc flasks was collected after 24 hours of continuous culture in selenium supplemented media at 100nM NaSe and 500nM SeMet. The collected cell pellet was resuspended in 300μL of cell lysis reagent (50mM Tris-HCl, 150mM NaCl, 1% Triton and 1% Tween-20) and incubated on ice for 30 min. The lysed cell suspension was centrifuged at 12000 rpm for 5 min to pellet cell debris and the protein-containing supernatant was removed and stored at -20°C until use.

**Protein Estimations:**
Quantification of the protein content from the cell extracts was performed using a BCA Protein Assay kit (Pierce, Rockford, USA) following the manufacturers protocol. Bovine serum albumin (BSA) was used as the protein standard, enabling the
construction of a standard curve, from which unknown protein concentrations could be extrapolated. Protein concentrations are expressed as mg of protein per mL of cell extract (mg/mL).

**Glutathione Peroxidase (GPx) Assay:**

Glutathione Peroxidase activity was quantified using a GPx assay, as first described by Flohe & Gunzler [9]. Glutathione Peroxidase present in the protein cell extract reduces the hydroperoxide substrate tert-butyl hydrogen peroxide and becomes oxidised. The oxidised GPx is regenerated by glutathione (GSH), which becomes oxidised to glutathione disulphide (GSSG). The GSSG can then be returned to GSH by the donation of two electrons from NADPH via the enzyme glutathione reductase (GR). The rate of decrease of NADPH directly correlates to GPx activity in the cell extract and can be measured spectrophotometrically at 340nm using a Tecan Sunrise plate reader with Magellan software. The activity measured was expressed as milliunits per milligram of protein (mU/mg of protein). The inter assay and intra assay coefficients of variation were 4.43% and 7.96% respectively.

**Thioredoxin Reductase (ThxRed) Assay:**

A commercial Thioredoxin Reductase assay was used to determine the activity in cells extracts. The kit was purchased from Cayman Chemical Company and 20 µL of cell extract was assayed in triplicate. The results were expressed as nmols activity per mg protein. The inter assay and intra assay coefficients of variation were 7.47% and 7.19% respectively.

**Selenium Supplementation and oxidative stress:**

Trophoblast cell number was adjusted to 100,000 cells per mL and 100µL was added to each well in a 96 well plate and incubated for 24 h at 37°C. Cells were then treated with selenium in the form of inorganic sodium selenite (NaSe) or organic selenium methionine (SeMet) prepared in media and added at concentrations of 100 nM or 500nM respectively. Cells were further incubated at 37°C for 24h. Trophoblast cells were then oxidatively stressed with various concentrations of Antimycin (40, 80, 160, 240, 340 µM) and Rotenone (100, 200, 400, 600, 800 nM) and incubated for 4 h at 37°C with 5% CO₂.
Resazurin End Point Assay:
To assess cellular function the Resazurin end point assay was used. This assay is
an oxidation-reduction indicator, detecting the production of NADH/NADPH and
assessing the mitochondrial metabolic activity [10]. Resazurin is non-fluorescent until
it is reduced to resorfin, which is highly fluorescent. Following treatments as
described above the media was removed by vacuum aspiration and cells were
washed twice with DPBS. Resazurin (200ul of 40nM) was added to each well
followed by a 2 hr incubation. The degree of reduction of resazurin to resorufin was
determined by fluorescence (excitation 530 nm; emission 590 nm) using a
Fluoroskan Ascent microplate fluorometer (Thermo Scientific, Australia).

Statistical Analysis
Statistical analysis was performed using Graph Pad, PRISM version 5.03 for
OneWay analysis of variance (ANOVA) with Tukey's Post Hoc testing was used to
analyse significant differences between selenium and non-selenium treatment
groups of equal n value for all treatment regimes. P values less than 0.05 were
considered significant and all data is presented as mean +/- standard deviation.
Each experiment was repeated a minimum of three times with all experiments
conducted in triplicate.

Results

Glutathione Peroxidase (GPx) and Thioredoxin Reductase (ThxRed) activity
post selenium supplementation
In BeWo cells, the baseline level of GPx activity was 129.6 ± 15.17 mU/mg (Fig 1).
After the supplementation with NaSe (100nM) or SeMet (500 nM), GPx activity was
increased to levels of 283.5 ± 14.24 mU/mg and 273.7 ± 14.49 mU/mg of protein. In
the BeWo cell line ThxRed baseline activity was 18.53 ± 2.11 mU/mg of protein. A
significant increase (P<0.05) in ThxRed activity was measured for cultures treated
with 100nM NaSe (72.14 ± 7.34 mU/mg of protein) and 500nM SeMet (69.52 ± 5.83
mU/mg of protein). Similar significant increases in GPx and ThxRed activity were
Selenium Supplementation Protects from Oxidative Stress induced by Rotenone treatment.

BeWo, JEG-3 and Swan-71 cell lines were cultured in either media only or supplemented with 100 nM NaSe or 500 nM SeMet to induce GPx and ThxRed activity. Cells were subsequently stressed with increasing doses of Rotenone (100, 200, 400, 800 nM) and cellular activity determined using the resazurin end point assay (Fig 2). Significant reductions in cellular activity were observed in cells exposed to all rotenone concentrations (P<0.001). Prior incubation with NaSe was demonstrated to protect trophoblast cells from oxidative stress at concentrations of 400 nM (70.99% ± 24.26; P<0.001) and 200 nM (88.56% ± 18.81; P<0.05) when compared to unsupplemented control cultures that displayed cellular activities of 36.77% ± 20.19 (400nM) and 71.72% ± 26.03 (200nM) respectively. Similarly, prior treatment with 500nM SeMet resulted in protection from rotenone-induced stress at concentrations of 400nM (66.92% ± 26.20; P<0.001) and 200nM (101.3% ± 15.96; P<0.05). Identical observations were made with JEG-3 and Swan-71 cells.

Selenium Supplementation Protection from Oxidative Stress induced by Antimycin treatment.

BeWo, JEG-3 and Swan-71 cell lines were cultured in either media or supplemented with 100 nM NaSe or 500 nM SeMet and subsequently stressed with increasing doses of Antimycin. Antimycin concentrations of 80µM to 320µM, applied for 4 hours, resulted in a dose dependent significant decrease in cellular activity in all three cell types (P<0.0001). As shown in Figure 3a, Swan-71 cells supplemented with NaSe exhibited protection from oxidative stress at Antimycin concentrations of 80µM (86.26% ± 16.45 vs. 52.60% ± 23.57; P<0.001), 120µM (75.06% ± 9.47 vs. 47.03% ± 12.09; P<0.01), 160µM (78.16% ± 21.72 vs. 26.74% ± 21.86; P<0.001) and 240µM (53.74% ± 22.88 vs. 11.51% ± 9.96; P<0.001). Prior treatment with SeMet also resulted in protection from Antimycin induced oxidative stress when exposed to concentrations of 80 µM (P<0.01), 120 µM (P<0.01), 160 µM (P<0.001) and 240µM (P<0.001) (Figure 3b). For BeWo and Jeg-3 cell lines, protection against Antimycin.
induced oxidative stress was observed only for the 320µM-160µM concentration range (P<0.001).

**DISCUSSION**

All aerobic tissues generate reactive oxygen and nitrogen species (RONS) and ample anti-oxidant production is essential for normal homeostasis. Oxidative stress arises when there is an excessive production of (RONS) or there is a diminished capacity of anti-oxidants to negate the detrimental effects of RONS. Typically a tissue experiencing oxidative stress will release evidence of this stress as oxidised lipids, proteins and nucleic acids. Excessive oxidative stress will result in tissue turnover and release of cellular components through apoptotic and necrotic mechanisms [1]. There are numerous reports on the release of oxidised macromolecules from the human placenta and oxidative stress has been proposed as a driver of cellular turnover in this tissue, especially during pathologies such as preeclampsia [11,12].

Glutathione peroxidase (GPx) and Thioredoxin reductase (ThxRed) are seleno-proteins and critical components of two very important anti-oxidant systems. Diminished GPx and ThxRed activity results in the generation of oxidative stress demonstrating a causative effect. Furthermore, there is decreased activity of these enzymes in tissues as a consequence of oxidative stress [4,7,11]. As seleno-proteins the expression and activity of GPx and ThxRed is dependent upon an adequate supply of selenium and the level of expression in cells and animal models can be controlled by selenium supplementation. Several reports have suggested that preeclamptic mothers are seleno-deficient and selenium intake may be important in the development of preeclampsia [4,13,14]. Hence, the purpose of this study was to investigate the link between selenium supplementation and protecting placental cells from oxidative stress.

Two forms of selenium, organic SeMet and inorganic NaSe were used to up regulate GPx and ThxRed in the trophoblastic cell lines BeWo, JEG-3 and Swan-71. Enzyme assays for GPx and ThxRed were used to confirm the up-regulation of activity post-supplementation. Similar observations have been made in many other cell types and in animal models [15,16]. The selenium dependent up-regulation of GPx and
ThxRed was most marked in BeWo and JEG-3 trophoblastic cell lines but less so in Swan-71 cells.

Following induction of anti-oxidant expression, the BeWo, JEG-3 and Swan-71 cells were oxidatively stressed with Rotenone and Antimycin in a dose dependent manner. Rotenone and Antimycin block Complex 1 and 3 respectively of the electron transport chain resulting in electron leakage and generation of superoxide (O\(^\bullet^-\) ) in the mitochondrial matrix. Due to the relatively high concentration and catalytic activity of Mn-SOD, the O\(^\bullet^-\) is rapidly converted to H\(_2\)O\(_2\) which is potentially very damaging to cell viability. In these experiments cellular metabolic activity was determined using the Reszaurin end point assay and we investigated whether up-regulation of GPx and ThxRed could protect from mitochondrial oxidative stress.

As presented above, there was a dose dependent decrease in the cellular activity in BeWo, JEG-3 and Swan-71 when treated with increasing concentrations of Rotenone (100-800nM) and Antimycin (40-320\(\mu\)M). Prior incubation with Na Selenite or Selenomethionine was able to protect trophoblast cells from oxidative stress. In all trophoblast cell lines there was considerable protection observed at Rotenone concentrations of 400-200nM. Selenium supplementation of Jeg-3 and BeWo cell lines conveyed protection at Antimycin concentrations 120-320\(\mu\)M, with protection against Antimycin concentrations as low as 80\(\mu\)M observed in Swan-71 cells. From these observations it would appear that there is a range of stressor concentrations where protection is possible and cells were irreversibly damaged by oxidative stress at higher concentrations. The mechanisms which cause this stress, whether it is lack of respiration, excessive RONS production, apoptosis or necrosis are currently subject to further investigation.

In a previous study we demonstrated that selenium supplementation could protect trophoblast cells from exogenously applied oxidative stress [8]. We used various forms of peroxides to stress cells and clearly demonstrated the beneficial effects of up-regulation of GPx and ThxRed through Se supplementation. However, the concentrations of peroxides used were probably higher than would be present in trophoblast cells in vivo and the application of exogenous peroxides would presumably have more of an effect in the cytoplasm as the cell absorbs the toxin. In
the present experiment we demonstrated a similar protective effect to endogenously generated oxidative stress. Blocking electron transport with Rotenone and Anyimycin is a generally accepted method of generating intra mitochondrial oxidative stress. This is very pertinent to studies on oxidative stress in the placenta during preeclampsia, which is often attributed to mitochondrial, and or endoplasmic reticulum stress. This study clearly shows a link between selenium, the antioxidant proteins GPx and ThxRed and mitochondrial oxidative stress.

Selenium is an essential micronutrient and without it tissues are susceptible to oxidative stress through loss of function of key enzymes such as GPx and ThxRed [13]. Selenium supplementation also offers an attractive method of maximising antioxidant expression in preparation of an oxidative insult [17]. Our previous work in cardiac tissues has shown the benefits of selenium supplementation in patients undergoing coronary by-pass surgery, a clinical scenario where oxidative stress negatively impacts on cardiac recovery [6]. Similarly, selenium supplementation may be of benefit for women who may experience preeclampsia. Indeed our recent work would suggest a link between selenium status and the development of preeclampsia and recent supplementation trials have indicated a possible beneficial effect [14,18]. Larger more controlled trials are needed to prove the benefits of selenium supplementation in lowering the incidence or severity of preeclampsia but it is now clear that selenium, through up-regulation of key anti-oxidant proteins can protect against mitochondrial oxidative stress in placental trophoblast cells.

REFERENCES.


**Figure 1:** Activity of GPx in NaSe and SeMet supplemented BeWo cell lines. Values are presented as mean ± standard deviation (SD). Significant differences were detected between control and maximum activity in the 100nM NaSe and 500nM SeMet supplemented cell lines (*p<0.05 n=3).
**Figure 2:** Activity of Thx-Red in NaSe and SeMet Supplemented BeWo and JEG-3 cell lines; quantified using Thx-Red inhibition assays. Values are presented as mean ± standard deviation (SD). Significant differences were detected between control and maximum activity in the 100nM NaSe and 500nM SeMet supplemented cell lines (*p<0.05, n=3).
**Figure 3.** Bewo trophoblast cells treated with NaSe (a) or SeMet (b) for 24hrs and subsequently treated with increasing concentrations of rotenone. Values are presented as mean ± standard deviation (SD) for each group. Significant differences were detected between unsupplemented and NaSe and SeMet supplemented cell lines when treated with 200nM and 400nM concentrations of rotenone (* p<0.05, ***p<0.001, n=9).
Figure 4: Swan-71 trophoblast cells treated with NaSe (A) or SeMet (B) for 24hrs and subsequently treated with increasing concentrations of antimycin. Values are presented as mean ± standard deviation (SD) for each group. Significant differences were detected between unsupplemented and NaSe and SeMet supplemented cell lines when treated with 80-240μM concentrations of antimycin (** p<0.01, ***p<0.001 n=9).