Increased circulating leukocyte numbers and altered macrophage phenotype correlate with the altered immune response to brain injury in metallothionein (MT) -I/II null mutant mice

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Abstract
Background: Metallothionein-I and -II (MT-I/II) is produced by reactive astrocytes in the injured brain and has been shown to have neuroprotective effects. The neuroprotective effects of MT-I/II can be replicated in vitro which suggests that MT-I/II may act directly on injured neurons. However, MT-I/II is also known to modulate the immune system and inflammatory processes mediated by the immune system can exacerbate brain injury. The present study tests the hypothesis that MT-I/II may have an indirect neuroprotective action via modulation of the immune system.

Methods: Wild type and MT-I/II−/− mice were administered cryolesion brain injury and the progression of brain injury was compared by immunohistochemistry and quantitative reverse-transcriptase PCR. The levels of circulating leukocytes in the two strains were compared by flow cytometry and plasma cytokines were assayed by immunoassay.

Results: Comparison of MT-I/II−/− mice with wild type controls following cryolesion brain injury revealed that the MT-I/II−/− mice only showed increased rates of neuron death after 7 days post-injury (DPI). This coincided with increases in numbers of T cells in the injury site, increased IL-2 levels in plasma and increased circulating leukocyte numbers in MT-I/II−/− mice which were only significant at 7 DPI relative to wild type mice. Examination of mRNA for the marker of alternatively activated macrophages, Ym1, revealed a decreased expression level in circulating monocyes and brain of MT-I/II−/− mice that was independent of brain injury.

Conclusions: These results contribute to the evidence that MT-I/II−/− mice have altered immune system function and provide a new hypothesis that this alteration is partly responsible for the differences observed in MT-I/II−/− mice after brain injury relative to wild type mice.

Keywords: Metallothionein, cryolesion, brain injury, alternatively activated macrophages

Background
Metallothionein (MT) is a 6-7 kDa, cysteine-rich, zinc-binding protein that has antioxidant properties. MT-I and MT-II are similar isomers, often considered to behave as one species (MT-I/II), that share the ability to provide protection to the injured brain. During insult to the central nervous system (CNS), metallothionein-I and -II double knockout (MT-I/II−/−) mice show increased neuron death or larger injury size after brain injury [1-3]. This firmly suggests that the presence of MT-I/II provides protection against CNS perturbation but the precise mechanisms that underlie this are yet to be identified. In vitro experiments have demonstrated that MT-I/II can provide protection, directly to neurons, against zinc toxicity [4] and can protect astrocytes from oxidative stress [5]. In a regenerative context, MT-I/II can enhance neurite extension in neurons [6]. However, a defining characteristic of brain injury in MT-I/II−/−
mice is the increased numbers of inflammatory cells such as microglia or macrophages, and T cells compared to wild type mice [2,7-8]. Notably, MT-1/I/II has been shown to affect immune system processes such as immunoglobulin production [9-14]. Leukocytes infiltrate the injured CNS and have the potential to be neurotoxic which makes it difficult to determine if the increased cell death observed in the injured brains of MT-1/I/II -/- mice is due to the absence of the neuroprotective effects of MT-1/I/II in the CNS or the absence of the modulatory effects that MT-1/I/II has on the immune system. The infiltration of neutrophils into CNS injuries is the most rapid of any type of leukocyte but neutrophils do not persist beyond 2 days post-injury, at which time monocytes become the dominant infiltrating leukocyte [15]. T cell infiltration occurs in several waves with an early infiltration within 1 hour [16], followed by a second infiltration at 24 hours [17]. However, the maximal T cell occupation of the injured CNS begins to occur about 1 week after the initial injury [18]. Evidence suggests that many immune system processes, such as inflammatory cytokine production and the oxidative burst, are neurotoxic and can impede the resolution of brain injury [19-21]. It is feasible that the number of immune cells entering the CNS can influence the progression of brain injury but the phenotype of the immune cells may also affect this process. For example, naive helper T cells can take on one of several phenotypes when they first become activated; the predominant types are type 1 helper T cell phenotype and the type 2 helper T cell phenotype [22]. Th1 cells promote the formation of classically activated macrophages (caMΦs) and augment the production of pro-inflammatory cytokines and reactive oxygen species and other neurotoxic molecules whereas Th2 cells promote formation of alternatively activated macrophages (aaMΦs) which antagonise these processes [23]. In vitro caMΦs have been shown to cause neuron death whereas aaMΦs appear to be less neurotoxic and possibly have some neuroprotective properties [24]. There is some evidence that T cells from MT-1/I/II -/- mice are more responsive to stimuli that induce Th1 cells [25] and differences in the numbers of circulating leukocytes in MT-1/I/II -/- mice relative to wild type mice have been observed [9]. Therefore, it is possible that the altered inflammatory response in the injured brain of MT-1/I/II -/- mice is a result of an altered immune system but this has not been investigated in depth.

In the present study we used a cryolesion injury model to compare the number of T cells infiltrating the injury site in wild type and MT-1/I/II -/- mice. Analysis of the numbers of leukocytes in circulation in the days following brain injury was also conducted to determine if the altered immune response to injury in MT-1/I/II -/- mice occurs before the cells enter the injured brain. Levels of the mRNA marker of aaMΦs, Ym1, were assayed to determine if MT-1/I/II -/- mice have different ratios of caMΦs/aaMΦs in comparison to wild type mice after brain injury.

Methods

Animals

All procedures involving animals were approved by the Animal Experimentation Ethics Committee of the University of Tasmania and were consistent with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes. Breeding stock for 129S1/SvImJ (wild type) mice and 129S7/SvEvBrd- Mtt1tm1Brt Mtt2tm1Brt/I (MT-1/I/II -/-) mice [26] were obtained from Jackson Laboratories. Male mice from both strains were housed with food and water ad libitum with 12/12 hour light/dark cycling. Mice used in the experiment were between 12 and 36 weeks of age. For each experiment, mice from both strains were divided evenly into groups of 5-7 animals for sampling time (0, 1, 3 and 7 days post-injury, DPI) and animals within these groups were randomised and placed in numbered cages to blind the strain of the mouse from the researchers. Each mouse was housed in an individual cage for at least 3 days prior to injury surgery and during the period after surgery until euthanasia.

Cryolesion brain injury

The cryolesion injury method was adapted from [27]. Mice were anaesthetised with 2-3% isoflurane/oxygen mix, delivered to the animal at 0.6 L/min. A 3 mm diameter, 50 mm long steel rod was cooled in liquid nitrogen. A sagittal incision along the skull was used to expose the cranium and the steel rod was then applied directly to the skull for 6 seconds. The stereotaxic coordinates for the injury site were 4 mm anterior of lambda and 2 mm right of the midline. The skin was sutured and the animal was allowed to recover back in its original cage. Mortality rate was less than 1% with a few animals showing signs of seizure within the first 24 hours after the application of the cryolesion injury. These animals were euthanased and excluded from the study.

Immunohistochemistry

Mice were transcardially perfused with phosphate buffered saline (PBS); brains were dissected out of the skull and drop-fixed in 4% paraformaldehyde for 24 hours. The brains were embedded in wax and sectioned at 10 μm thickness. Before staining, antigen retrieval was undertaken in 0.01 M citrate buffer, pH 6, in a pressure cooker for 10 minutes. Primary antibodies used were 1:100 rat monoclonal NIMP-R14 to neutrophil (Abcam, Cambridge, UK) for neutrophils, 1:500 goat polyclonal
anti-Iba1 (Abcam) for microglial/mixed cell derived macrophages and 1:100 rabbit polyclonal anti-CD3 (Abcam) was used for T cells. All antibodies were diluted with 0.3% Triton-X 100 (Sigma, St. Louis, MO, USA) solution in PBS. Blocking with serum-free protein block (Dako, Glostrup, Denmark) was required for CD3 staining and was applied for 30 minutes before application of the primary antibody. The diluted NIMP-R14 antibody solution contained 10% normal goat serum (Vector Labs, Burlingame, CA, USA) as a blocking reagent. Biotinylated goat anti-rat IgG (Invitrogen, Carlsbad, CA, USA), biotinylated goat anti-rabbit IgG (Invitrogen) or donkey anti-goat IgG (Santa Cruz, Santa Cruz, CA USA) secondary antibodies, were applied to sections at 1:1000 dilution for 1 hour at room temperature. Avidin-biotin complex (Vector Labs) was used as the detection reagent and was applied to sections for 15 minutes followed by 2 rinses in PBS. Nickel enhanced 3’-diaminobenzidine (DAB, Vector Labs) was used as the chromogen and was applied at the manufacturer’s specified concentration for 8 minutes. Slides were then rinsed in distilled water for at least 5 minutes. Nuclear Fast Red (Sigma) was used as a counterstain for NIMP-R14 and Iba1 stained sections.

Fluoro-jade C staining
Fluoro-jade C (Millipore, Billerica, MA, USA) is a neuron-specific marker of dead and degenerating neurons. Staining was carried out according to the protocol of Schmued et al. [28] whom demonstrated that fluoro-jade C labels both apoptotic and necrotic neuronal death without discrimination. Rehydrated, slide-mounted sections were immersed in 0.06% potassium permanganate solution for 10 minutes. The slides were rinsed for 2 minutes in distilled water then immersed in 0.001% fluoro-jade C, 0.01% acetic acid solution for 10 minutes. The slides were rinsed twice in distilled water for 5 minutes then were air-dried before they were coverslipped with DPX mounting medium (Merck, Whitehouse Station, NJ, USA).

In situ cell counts in the injured brain
Low power, 10 x objective images were taken of the injury site for sections stained for fluoro-jade C, NIMP-R14, Iba1 and CD3. Fluoro-jade C counts were conducted for all positively labelled cells in the injury site and at lower depths in the un-injured CNS parenchyma. To standardise cell counts, fluoro-jade C counts were divided by the linear width of the injury in the section plane at the upper cortical surface in millimeters. NIMP-R14, Iba1 and CD3 positive cells were only counted within the injury site. Cell counts within the injury site were standardised to the area of the injury site in that section in mm². The injury border was demarcated by an obvious degradation of tissue integrity observed in the injury site. The border of this region correlated well with the GFAP⁺ endfeet extended by astrocytes as they re-established the glia limitans at days 3 and 7 DPI (data not shown). The glia limitans was not re-established at this injury border at 1 DPI but the pyknotic nuclei stained by nuclear fast red that likely represent apoptotic cells were rarely found outside this zone of reduced tissue integrity. Therefore this boundary was deemed to be a physiologically relevant demarcation of the injury zone. All cell counts were conducted blinded to the strain of the mouse.

Flow cytometry
Blood was obtained by cardiac puncture and the leukocytes were separated from erythrocytes by Histopaque 1119 (Sigma) density gradient centrifugation. For CD3 and CD4 double labelling, 10⁶ leukocytes were used for each batch of staining. Leukocytes were stained with a combination of 1 μg/ml APC-conjugated hamster IgG1 anti-mouse CD3e (BD Biosciences, Franklin Lakes, NJ, USA) and 1 μg/ml PE-conjugated rat IgG2a anti-mouse CD4 (BD biosciences) in 200 μl PBS-2%FCS at 4°C for 15 minutes. The cells were pelleted by centrifugation, and washed twice by resuspension in PBS-2%FCS followed by centrifugation to pellet. The pellet was resuspended in a fixation solution consisting of 2% paraformaldehyde, 4% D-glucose, 0.03% sodium azide and 0.01 M PBS for storage. For CD4⁺CD25⁻FoxP3⁺ naturally occurring regulatory T cells, 10⁶ cells were used for each batch of staining and were labelled with the mouse regulatory T cell staining kit # 2 (eBioscience, San Diego, CA, USA) according to the manufacturer’s protocol. Staining procedures were also carried out for isotype control antibodies, PE-conjugated rat IgG2a (BD Biosciences) and APC-conjugated hamster IgG1 (BD Biosciences), which were applied at the same concentration as the specific antibodies to unstained cells. Samples were assayed by flow cytometry (BD Canto II flow cytometer) and were analysed using BD FACS Diva software v6.1.1. A quadratic gate was applied to the scatter plots of CD3 versus CD4 fluorescence to identify CD3⁺CD4⁺ and CD3⁻CD4⁺ T cells which were expressed as a percentage of all leukocytes. To identify naturally occurring regulatory T cell populations, a gate was applied to cells expressing CD4. Cells within the CD4⁺ population gate were analysed with a quadratic gate applied to the scatter plots of CD25 versus FoxP3. CD4⁺CD25⁻FoxP3⁺ cells are expressed as a percentage of CD4⁺ cells. The distinction between positive and negative staining was determined by the upper fluorescence of isotype control stained cells. All thresholds and gates were applied on this basis. To determine absolute circulating leukocyte numbers, blood was obtained from mice by cardiac puncture with syringes containing...
EDTA (3 mg per ml of blood). From each animal 250 μl whole blood was analysed in an Advia 120 haemocytological analyser (Siemens, Munich, Germany).

Quantitative reverse-transcriptase PCR (RT-PCR)
Mice were transcardially perfused with PBS. The brain injury site was dissected out of the brain using a 3 mm biopsy punch and homogenised via Ultra-Turrax (IKA, Staufen, Germany) in TRI-reagent (Sigma, St. Louis, MO, USA). Peripheral blood mononuclear cells (PBMCs) were obtained using density gradient centrifugation on Histopaque 1083 (Sigma) and the pelleted cells from this fraction were homogenised in TRI-reagent. RNA was isolated from TRI-reagent according to the manufacturer’s protocol. Reverse transcription with the Superscript-III reverse transcriptase system (Invitrogen) and quantitative PCR with Quantitect SYBR green (Qiagen, Hilden, Germany) was conducted according to the method of Brettingham-Moore et al. [29]. Oligonucleotide primers are detailed in table 1. The MT-I and MT-II primer sets were designed to be complementary to the cDNA for the transcripts from both wild type and MT-1/II−/− mice, which still produce MT-I and MT-II transcripts but have premature stop codons inserted to prevent complete protein translation. Standard curves were created using known quantities of each PCR product and were used to determine the original cDNA copy number at an arbitrary fluorescence threshold (C_T). GAPDH mRNA was used as the housekeeping gene and MT-I and MT-II mRNA copy numbers were standardized to the copy number of the house-keeping gene, GAPDH.

Plasma cytokine assay
Blood was collected from mice via cardiac puncture with heparinised syringes. Plasma was obtained after centrifugation of blood for 5 minutes at 14000g. Plasma samples were diluted four-fold with PBS and assayed with a cytometric bead array mouse Th1/Th2/Th17 cytokine kit (BD biosciences). The assay was run according to specification and analysis was conducted using FCAP Array software v1.0.1. IL-4 and IFN-γ were assayed with ELISA kits (R&D systems, Minneapolis, MN, USA) according to manufacturer’s protocol.

Statistical Analysis
Statistical analysis was conducted with SPSS 15.0 (SPSS Inc.). Homogeneity of variances between groups within each data set was determined with Levene’s test. Box-Cox test was used to determine the appropriate transformation for data sets with heterogeneous variances between groups. Statistical significance was determined with two-way ANOVA for p-values < 0.05 with Tukey’s B Post-hoc test. All error bars in figures represent the standard error of the mean (SEM).

Results
Histological comparison of the extent of injury in wild type and MT-1/II−/− mice
The area of the injury site in a 5 μm section taken from the widest point of the injury site was used as a comparative measure of injury size. The size of the injury did not change significantly from 1-3 DPI but declined significantly from 3-7 DPI in both wild type mice and MT-1/II−/− mice (Figure 1A). No significant differences were observed between the strains at any time-point which suggests that on a large scale, the severity of the injury and rate of healing is similar in wild type and MT-1/II−/− mice. However, investigation of the death of individual neurons revealed some notable differences between the two strains of mouse. Fluoro-jade C is a histological dye that labels neurons dying by both apoptosis and necrosis [28] and was found to label neurons in the lesion site and in the surrounding, uninjured parenchyma (Figure 1B). Quantification of fluoro-jade C labelled cells revealed that the highest degree of cell death occurred at 1 day post-injury (DPI, Figure 1B). In wild type mice the number of fluoro-jade C labelled cells decreased from 1-3 DPI and again at 7-3 DPI. MT-1/II−/− mice had a similar decrease in fluoro-jade C staining from 3-1 DPI compared to wild type mice. In contrast to wild-type mice, the amount of cell death did not differ between 3 and 7 DPI in the injury site of MT-1/II−/− mice. As a result, there were significantly greater numbers of fluoro-jade C labelled cells in MT-1/II−/− mice at 7 DPI than in wild type mice.

Leukocyte infiltration into the injury site in wild type and MT-1/II−/− mice
Leukocyte infiltration into the cryosections at 1, 3 and 7 DPI was investigated to determine if leukocytes could

| Table 1 Oligonucleotide primer sets used for quantitative RT-PCR of brain mRNA samples after cryoleision brain injury. |
| --- | --- | --- |
| Primer | Forward (5' - 3') | Accession No. |
| GAPDH | CCCAGAAAAGACGTTGGATG | NM_008064.2 |
| Rev | GGATCGAGGATGTCT | |
| IFN-γ | ACTGCCAAGAAGATGGTGC | NM_008337.3 |
| Rev | GACCTGGCTCGATGACCT | |
| IL4 | CACACCCGGCAGCTTGTCC | NM_021283.2 |
| Rev | TCTTGGCAGTCTGTC | |
| Ym1 | ACAATTTAGAGGGTCCTGG | NM_009652.2 |
| Rev | CCAGCTGAGTCTCCTGG | |
| MT4 | GCCGTCACCTACAGCTG | NM_013602.3 |
| Rev | AGCACGACAGCTTGG | |
| MT-II | CAAAGCAGGATCCCTGG | NM_008330.2 |
| Rev | AGCACGACAGCTTGG | |
Figure 1 Quantification of injury size and neuron death after cryolesion injury in wild type (grey bars) and MT-III<sup>−/−</sup> mice (white bars). Injury size (A) was quantified by measurement of the area of the injury in sections taken from the widest point of the injury site. Neuron death identified by fluor-jade C labelling (B). Fluor-jade C<sup></sup>-positive cells were counted in the injury site and the surrounding tissue. Counts were standardized per linear mm (width) of the injury site. Lower case letters indicate signficance determined by Tukey's B post-hoc test. Time-points sharing letters indicates lack of statistically significant difference. n = 5–7, error bars = SEM.

Representative images of fluor-jade C staining in the injury site of wild type animals at 1 DPI (C), 3 DPI (D) and 7 DPI (E) with scale bars = 200 μm.

play a role in the sustained neuron death at 7 DPI in MT-I/II<sup>−/−</sup> mice. Neutrophils were identified by immunohistochemistry for NIMP-14 and were at their highest levels in the injury site at 1 DPI coinciding with the highest amount of neuron death (Figure 2A).

Neutrophil numbers were greatly diminished at 3 DPI and mostly absent from the injury site at 7 DPI. No significant differences were found between neutrophil numbers in the injury site of wild-type and MT-I/II<sup>−/−</sup> mice at 1, 3 or 7 DPI. Neutrophils were found mainly within the injury site but were occasionally found in the uninjured parenchyma proximal to the injury border.

Iba1 staining was used to identify macrophages derived from activated microglia and infiltrating monocytes within the injury site (Figure 2B). Macrophages within the injury site increased from 1–3 DPI and reached maximal numbers for the study period at 7 DPI. No significant differences in macrophage numbers between MT-I/II<sup>−/−</sup> and wild type mice were observed at any time point.

CD3 antibody was used to identify T cells in the injury site (Figure 2C). T cells were found to be mainly confined to the injury site. T cell numbers were found to be relatively low at 1 and 3 DPI with no significant differences between injuries of wild type and MT-I/II<sup>−/−</sup> mice. However, at 7 DPI, T cell numbers were greatly increased compared to the earlier time points. MT-I/II<sup>−/−</sup> mice had significantly more T cells per mm<sup>2</sup> in the injury site than wild type mice at 7 DPI.

**Metallothionein expression in the cryolesion injury site**

The levels of MT-I and MT-II mRNA were assessed post-injury in wild type mice by quantitative RT-PCR. MT-I (Figure 3A) and MT-II (Figure 3B) mRNA both show significant increases in expression relative to GAPDH at 1 DPI. MT-II appears to be the dominant isoform of MT with 41.1 fold higher levels of expression than MT-I relative to GAPDH. At 3 and 7 DPI, MT-II mRNA was decreased but remained significantly elevated relative to the uninjured cortex, whereas MT-I mRNA had returned to pre-injury levels. It is interesting to note that peak MT-I and MT-II mRNA expression in the brain does not coincide with observed differences in neuron death and T cell infiltration in wild type and MT-I/II<sup>−/−</sup> mice.

**Circulating leukocyte numbers in wild type and MT-I/II<sup>−/−</sup> mice after brain injury**

Previous studies have found MT-I/II<sup>−/−</sup> mice to have altered levels of circulating leukocytes and leucocyte subtypes [9]. A haematological analyser was used to determine whether differences in absolute numbers of white blood cells in MT-I/II<sup>−/−</sup> mice and wild type mice might account for differences in leucocyte infiltration rates into the cryolesion-affected tissue (Figure 4A). Leukocyte counts from whole peripheral blood did not significantly change after brain injury in wild type mice at 0 (uninjured), 1, 3 or 7 DPI. MT-I/II<sup>−/−</sup> mice had no significant changes in leucocyte numbers in whole peripheral blood from 0-3 DPI but had significantly higher leucocyte counts at 7 DPI compared to uninjured controls and wild type mice at 7 DPI (Figure 4A). Analysis of circulating neutrophils, lymphocytes and monocytes revealed no differences in the relative ratios of any leucocyte sub-type (Figure 4B). Basophils and eosinophils constituted a small fraction of all leucocytes and numbers did not increase after injury in MT-I/II<sup>−/−</sup> mice and do not explain the increased levels of leucocytes in MT-I/II<sup>−/−</sup> mice at 7 DPI (data not shown). Because the haematological analyser did not differentiate between sub-classes of lymphocytes, flow cytometry was used to determine if relative ratios of T cells were equal in MT-I/II<sup>−/−</sup> mice and wild type mice (Figure 5A). By 7 DPI, there was a significant overall decrease in CD3<sup></sup> CD4<sup></sup> helper T cells in both MT-I/II<sup>−/−</sup> mice and wild type mice. However, there was no significant difference at any timepoint between these two groups of mice. There was also no difference in numbers of CD3<sup></sup> CD4<sup></sup> T cells, the majority of which are likely to consist of cytotoxic T cells (data not shown).
CD4+CD25+FoxP3+ naturally occurring regulatory T cells have been shown to reduce the impact of stroke [30] and may have similar protective roles in the injured brain. However, the number of CD4+CD25+FoxP3+ naturally occurring regulatory T cells as a percentage of CD4+ T cells was not found to vary significantly at 3 or 7 DPI between wild type and MT-I/II+ mice (Figure 5C). Therefore, there were no differences in the ratios of any of the leukocyte sub-types investigated in wild type and MT-I/II+ mice after brain injury. However, the increase in absolute leukocyte numbers observed in MT-I/II+ mice at 7 DPI allows us to calculate that there would be an average increase of 16% in the absolute number of circulating T cells in MT-I/II+ mice at 7 DPI. Because there was no increase in the absolute number of leukocytes in wild type mice, the same calculation determines that absolute numbers of circulating T cells would have decreased by 26% on average by 7 DPI. This result is in accordance with the finding that T cell numbers are lower in the brain of wild type mice compared to MT-I/II+ mice at 7 DPI.

Comparison of chemokine and cytokine expression in wild type and MT-I/II mice

Plasma concentrations of Th1 and Th2 cytokines were assayed to determine if there were systemic differences in the inflammatory response to brain injury. Unfortunately most of the cytokines tested were not detectable or were only rarely detected (IL-4, IL-6, INF-γ, TNF-α and IL-10) which suggests that the cryopreservation does not induce a strong systemic inflammatory response. However, it is known that T cells must be activated before they can enter the CNS [31,27] and the cytokine, IL-2 which is responsible for T cell activation, was detectable
in the plasma of some animals after injury (Figure 6). At 1 and 3 DPI, IL-2 was detected in some animals from both the wild type and MT-1/II−/− groups. At 7 DPI, only MT-1/II−/− mice had detectable plasma concentrations of IL-2 with 4 out of 6 animals exhibiting detectable expression of IL-2. No wild type animals had detectable levels of IL-2 at 7 DPI. Statistical analysis could not be conducted on this data set due to the high number of animals with plasma IL-2 concentrations lower than the detection limit.

IL-4 and IFN-γ mRNA could not be detected in the injury site of either strain of mouse, yet both transcripts were detectable in RNA harvested from a mouse T cell line that had been stimulated with calcium ionophore and phorbol ester to induce a state of activation (data not shown). IL-4 and IFN-γ protein could not be detected in the injury site of MT-1/II−/− mice and wild type mice. Overall, systemic cytokine activation was not observed after brain injury and local T cell specific cytokines were undetectable so Th1 and Th2 responses could not be compared directly in wild type and MT-1/II−/− mouse injury sites.

To assess the effect of increased numbers of T cells in the injury site, quantitative RT-PCR was used to assess the levels of mRNA for the alternative macrophage activation marker, Ym1, in the cryolesion site (Figure 7A). Ym1 expression increased significantly at 1 DPI in the injury site of wild type and MT-1/II−/− mice as determined by 2-way ANOVA. It was also determined that wild type mice have significantly higher levels of Ym1 compared to MT-1/II−/− mice independent of the factor of time before or after injury.

Using this method, it was impossible to determine whether the Ym1 is derived from the CNS-resident microglia or infiltrating monocytes because both cell types contribute to the pool of activated macrophages in the injury site. To examine whether the increased Ym1 in wild type mice occurs in monocytes before they enter the injured brain, RT-PCR was used to determine Ym1 levels in PBMCs (Figure 7B). Monocytes are the only cell type in this cell fraction that express Ym1. 2-way ANOVA revealed that there were no significant changes in Ym1 expression over time, but wild type animals express significantly higher levels of Ym1 mRNA in their PBMCs compared to MT-1/II−/− mice.
Discussion

The present study demonstrates that the altered immune response present in MT-I/II−/− mice occurs in later stages of brain injury. The significant findings of increased T cell infiltrate into the injury site, increased levels of IL-2, prolonged neuronal death surrounding the injury site and increased numbers of circulating leukocytes were all only present in MT-I/II−/− mice at 7 DPI. However, the finding that MT-I/II−/− mice have altered expression of Ym1 mRNA in the brain and circulating monocytes, both before and after injury, suggests that at least some aspects of the altered immune response in MT-I/II−/− mice are independent of brain injury. The implication of these findings is that MT-I/II may be having effects on the immune system systemically hence MT-I/II may be affecting the progression of brain injury indirectly via modulation of inflammatory responses.

A latent period before increased neuron death in MT-I/II−/− mice compared to wild type mice has been reported previously [1] which is in accordance with the findings of the present study. However, increased neuron death in MT-I/II−/− mice has also been observed within 24 hours of brain injury by Penkowa et al. [2]. One of the criticisms of the study by Penkowa et al. [2] is that neuron death in the cryolesioned cortex was assessed by counting the number of remaining neuron-specific enolase labelled neurons immediately adjacent...
Figure 6 Scatter plot showing detectable plasma IL-2 concentrations in MT-I/II<sup>−/−</sup> mice (crosses) and wild type mice (circles) after brain injury. Values below the detection limit (0.4 pg/ml) are not shown. Increases in IL-2 in plasma after injury were spurious with few animals posting detectable concentrations. At 7 DPI only MT-I/II<sup>−/−</sup> mice have detectable levels of plasma IL-2. n = 7 for all groups except wild type mice at zero DPI and MT-I/II<sup>−/−</sup> mice at 7 DPI for which n = 6, error bars = SEM.

to the injury border. However in that study, the size of injury site was larger in MT-I/II<sup>−/−</sup> mice compared to the wild type controls, hence neuron counts from MT-I/II<sup>−/−</sup> mice would have come from deeper cortical layers than in wild type mice. It is well known that there are large differences in the density and distribution of neurons in different layers of the cortex [32]. Natale et al. [1] also used counts of remaining neurons to measure neuron death in brain injury but their method of brain injury allowed counting to be conducted in the same region in both wild type MT-I/II<sup>−/−</sup> mice. In the present study there were no significant differences between injury sizes in wild type and MT-I/II<sup>−/−</sup> mice and cell counts were conducted based on a specific marker of neuronal death.

Crowthers et al. [9] have reported that MT-I/II<sup>−/−</sup> mice have altered numbers of T cells in the blood and spleen when compared to wild type mice but other investigators found no difference in T cell numbers in the spleen and lymph nodes of wild type and MT-I/II<sup>−/−</sup> mice [25]. In the present study, leukocyte numbers in MT-I/II<sup>−/−</sup> mice were only found to differ after brain injury. Extracellular MT-I/II has been shown to have modulatory effects on multiple types of leukocytes [9,14,33,34]. Increases in extracellular MT have been observed in the blood of head injured patients despite the fact that MT-I/II protein has no secretory signal sequence [35]. However, the concentration of MT-I/II required to modulate the activity of immune cells in vitro is often higher than the levels of MT observed in circulation before or after head injury so further study is required to determine whether this mechanism is possible under the physiological conditions that occur after brain injury.

A possible mechanism by which MT-I/II acts on the immune system that has received little attention is that the zinc-binding ability of MT-I/II may affect immune system functioning. Zinc supplementation in humans has been shown to enhance leukocyte responses to activating stimuli [36] and zinc homeostasis is known to be disrupted by brain injury [37]. We have recently demonstrated that zinc is released from hepatic stores in mice after brain injury and that MT-I/II<sup>−/−</sup> mice have a reduced capacity for zinc sequestration to the liver, a process that occurs at 7 DPI in wild type mice (Pankhurst et al., manuscript submitted). The co-occurrence of this event with many of the altered immune system responses observed at 7 DPI in the present study provides evidence that MT-I/II mediated zinc homeostasis may be linked to immune system functioning.

We can not exclude the possibility that MT-I/II is interacting with more than one process that affects the injured brain. Increased Ym1 mRNA expression is a marker for aAMs [38] and was found to be significantly higher in wild type mice than MT-I/II<sup>−/−</sup> mice in both the injury site and PBMCs. The fact that Ym1 was higher in wild type mice compared to MT-I/II<sup>−/−</sup> mice before injury implies that wild type macrophages have a greater intrinsic disposition to become aAMs than those from MT-I/II<sup>−/−</sup> mice. This intriguing observation

Figure 7 (A) Ym1 mRNA expression is greater in the injury site of wild type mice (solid lines) than in MT-I/II<sup>−/−</sup> mice (dashed lines), n = 6-7, error bars = SEM. (B) Ym1 mRNA expression is significantly greater in the circulating PBMCs of wild type mice (solid lines) than in MT-I/II<sup>−/−</sup> mice (dashed lines), independent of time after injury, n = 6, error bars = SEM.
is likely to be independent of effects on zinc homeostasis in the liver of mice lacking MT-1/II. This trend was retained throughout the period after brain injury and may be partly responsible for the increased neuron death that was observed at 7 DPI in MT-1/II-/- mice compared to wild type mice. In vitro, the aaMφ response has been shown to be much less neurotoxic than the caMφ response which is purported to be due to the higher production of reactive oxygen species by caMφs [24]. CaMφs also produce higher levels of neurotoxic metabolites via the quinolinic acid pathway [39,40]. Th1 cytokines are responsible for the generation of caMφs and Th2 cytokines are responsible for the generation of aaMφs. We have previously shown that exogenous application of MT-1/II to the injured rat brain leads to a reduction in quinolinic acid production and extracellular application of MT-1/II to cultured microglia reduces Th1 cytokine-mediated (IFN-γ) production of quinolinic acid [41]. This is supported by the finding that naive T cells isolated from MT-1/II-/- mice have been shown to be more responsive to becoming Th1 cells than T cells from wild type mice [25]. One limitation of the present study is that measurement of the Th1/Th2 responses were not possible and we have only provided a single marker of caMφs hence more experiments are required to determine if differential macrophage activation is occurring in MT-1/II-/- mice and whether Th1/Th2 ratios differ. Immunoblot and RT-PCR of cytokines were not sensitive enough to definitively determine the relative ratios of the Th1 and Th2 responses in the injured brain in the present study. This was most likely due to the small tissue sample sizes and the fact that cytokines can operate at very low concentrations. However, IL-2 was detectable in the plasma of some animals and it was interesting to find that at 7 DPI only MT-1/II-/- mice were producing detectable amounts of IL-2. IL-2 is responsible for the clonal expansion of activated T cells [42] and we regard this as evidence that T cell activity was altered in MT-1/II-/- mice after brain injury and may explain the differences observed in T cell infiltration in MT-1/II-/- mice.

Conclusions
MT-1/II expression is increased in the brain after brain injury which suggests that some of the protective effects of MT-1/II after brain injury are acting directly on the injured brain. However, many of the processes observed in the current study are initiated outside the CNS. Therefore, it is possible that MT-1/II produced outside the injured brain could be more important for the modulation of immune response after brain injury than MT-1/II produced within the CNS. Such an interaction would increase the prospects for the use of MT-1/II as a therapeutic for brain injury.

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Authors’ contributions
MWP conceived the experimental design and conducted the majority of the experimental procedures and statistical analysis. UJt developed the immunohistological techniques for the study and participated in experimental procedures. ARW, SATK and RSC were involved in the development of the experimental approach and contributed significantly to interpretation of the data and preparation of the manuscript. All authors have read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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