Changes in Subtypes of Ca Microdomains Following Partial Injury to the Central Nervous System

Ivan Lozića,b, Carole A. Bartlettb, Jeremy A. Shawc, K. Swaminathan Iyera, Sarah A. Dunlopb, Matt R. Kilburnia and Melinda Fitzgeraldb*

Rapid changes in Ca2+ concentration and location in response to injury play key roles in a range of biological systems. However, quantitative analysis of changes in size and distribution of Ca2+ microdomains in specific cell types in whole tissue samples has been limited by analytical resolution and reliance on indirect Ca2+ indicator systems. Here, we combine the unique advantages of nanoscale secondary ion mass spectrometry (NanoSIMS) with immunohistochemistry to directly quantify changes in number, size and intensity of Ca microdomains specific to axonal or glial regions vulnerable to spreading damage following neurotrauma. Furthermore, using NanoSIMS allows separate quantification of Ca microdomains according to their co-localization with areas enriched in P. We rapidly excise and cryopreserve optic nerve segments from adult rat at time points ranging from 5 minutes to 3 months after injury, allowing assessment of Ca microdomains dynamics with minimal disruption due to tissue processing. We demonstrate significantly more non-P co-localized Ca microdomains in glial than axonal regions in normal optic nerve. The density of Ca microdomains not co-localized with P rapidly, selectively and significantly decreases after injury; densities of Ca microdomains co-localized with P enriched areas are unchanged. An efflux of Ca2+ from microdomains not co-localized with P may contribute to the structural and functional deficits observed in nerve vulnerable to spreading damage following neurotrauma. NanoSIMS analyses of Ca microdomains allow quantitative and novel insights into Ca dynamics, applicable to a range of normal, as well as diseased or injured mammalian systems.

Introduction

Rapid alterations in Ca2+ concentration and location are key features of the injury response in a range of biological systems. Injury to the central nervous system (CNS) results in damage to neurons and glia, with rapid alterations in Ca2+ concentration thought to be an important contributor1-4. Undamaged tissue adjacent to the primary injury is also susceptible to cell death through a series of disruptive cellular and metabolic cascades, termed secondary degeneration5. Secondary degeneration is thought to be triggered by glutamate-induced excitotoxicity, resulting in excess influx of Ca2+ into cells and associated oxidative stress6-8. While fluctuations in Ca2+ are a normal and essential part of neural function, excess Ca2+ influx into cells is a feature of CNS injury and is thought to contribute to the death of neurons and glia vulnerable to secondary degeneration1,4. The dynamic changes in Ca2+ flux in tissue vulnerable to secondary degeneration have not been fully characterized in vivo. Nevertheless, it has been proposed that movement of Ca2+ between glia and neurons could play a part in the spread of Ca2+ from the primary injury site, to areas affected by secondary degeneration9,10.

Ca2+ has been observed throughout the cytosol2, and in in vitro studies has also been shown to be localized in areas of increased concentration, referred to as microdomains11. Microdomains of Ca2+ are associated with presynaptic Ca2+ channels, as well as other cell features in the soma, including the mitochondria, Golgi body and nucleus11-13. Ca2+ microdomains are involved in synaptic activation, AMP signaling and presynaptic transmitter release14 and are found in astrocyte somata and processes13,15. In vivo and ex vivo 2-photon imaging has been used to demonstrate Ca2+ signaling using indicator dyes and genetically encoded Ca2+ indicators in normal and injured white matter tract16,17, including in astrocytes13,18-20, with the advantage of real-time recording21. However, the spatial resolution of the resultant images is limited. Quantification of changes in Ca2+ signal associated with microdomains in individual cellular compartments has been confined to small transients in sub-regions of 4 - 12 μm referred to as ‘focal’ events19, and Ca signals specifically associated with individual microdomains are yet to be comprehensively characterized22. Furthermore, these techniques rely on indirect measures of Ca2+, detecting changes in the indicator, rather than changes in Ca2+ itself.

Here, we combine nanoscale secondary ion mass spectrometry (NanoSIMS) with immunohistochemistry to characterize Ca microdomains in normal optic nerve, quantitatively investigating changes in Ca microdomains specific to axons or glia vulnerable to secondary degeneration, in the 3 months after partial injury. NanoSIMS is a high resolution imaging mass spectrometry technique that uses a finely focused ion beam to extract the top few atomic layers of a sample surface, producing an image reflecting the elemental/isotopic composition of the sample. It is...
highly versatile, and has been used to investigate cell structure and map the distribution of stable isotopes, metal species, and organometallic complexes within biological tissue. Furthermore, NanoSIMS has the important advantage of directly measuring the abundance of ions such as Ca, rather than relying on indirect measures of changes in indicator dyes. In a preliminary study, we used NanoSIMS to directly visualize the redistribution of areas of elevated Ca in optic nerve vulnerable to secondary degeneration within the first day of partial optic nerve transection injury.

In the current study we demonstrate that this high-resolution technique can also be used to quantify changes in Ca signals associated with individual microdomains as small as 1 μm in diameter, specific to glial or neuronal regions identified immunohistochemically. We define Ca microdomains as localised areas of increased Ca concentration likely to be associated with a Ca$^{2+}$ signal, which can merge together to form larger microdomains or macrodomains. The Ca microdomains we describe in the current study are distinct from Ca nanodomains, which are highly localized, transient phenomena, smaller than 1 μm and associated with a single ion channel. It is important to note that our cryopreservation protocol does not include fixation, and as such, it is possible that transient Ca microdomains were captured at the moment of high-pressure freezing. We assess fields of view from central optic nerve not damaged by the initial partial injury and shown to be vulnerable to secondary degeneration following dorsal partial transection. We examine ex vivo central optic nerve regions by rapid excision and cryopreservation of tissue, thereby ensuring that, as far as possible, cellular activity is immobilized such that the tissue closely approximates its living state. A further novel advantage of our current application of NanoSIMS is that Ca microdomains can be categorized and quantified on the basis of their co-localization with nano-scale areas enriched in Phosphorus (P) and therefore with areas that are not likely to have a higher concentration of phosphorylated proteins. Receptor mediated events involving Ca release can be either dependent or independent of phosphorylation. Changes in Ca$^{2+}$ microdomains differentiated according to co-localization with areas enriched in P, may thus implicate changes in functionally important regulatory mechanisms in specific cell types vulnerable to secondary degeneration.

**Experimental**

**Animals and Partial Optic Nerve Transection**

Adult, female piebald viral Glaxo (PVG) black hooded rats (165-205 g, Animal Resource Centre, Murdoch, WA, Australia) were used as described. Procedures were approved by The University of Western Australia's Animal Ethics Committee and conformed to the National Health and Medical Research Council of Australia guidelines on the use of animals in research. Partial optic nerve transection of the right optic nerve was performed as described previously.

**Sample Preparation for NanoSIMS**

Optic nerve tissue segments were collected 5 min, 1, 3, 7 days, and 3 month post partial optic nerve transection (n = 3 - 4 / time point). Segments (200 μm) of right optic nerve from uninjured animals were used as controls (n = 3). Transverse segments of the right optic nerve encompassing the injury site (200 μm) were rapidly excised using a custom-built cutter incorporating parallel razor blades placed 200 μm apart. The process of excision, cryoprotection and freezing was completed within 90 seconds. NanoSIMS analyses are similar to ultrastructural studies in that, due to practical considerations, only a relatively small number of samples can be assessed. Numbers of animals used per group for this new application of NanoSIMS technology was based on previously published ultrastructural analyses; blinding of investigators to sample identity was not possible. Subsequent to excision, each segment was briefly dipped in cryoprotectant solution (2% low-temperature gelling agarose, Sigma, St. Louis, MO), transferred to a membrane carrier (1.5 x 0.2 mm diameter, Leica) and high-pressure frozen (EMPACT 2, Leica) using liquid N$_2$, enabling a freezing rate of approximately 25,000 °C/sec with minimal ice crystal formation. A protocol outlined by Clode and Marshall was used to minimize movement of mobile ions such as Ca. In brief, frozen optic nerve samples were freeze-substituted (EM AF52, Leica) in 10% acrolein in Et$_2$O and tissue was infiltrated with Lowicryl HM20 resin (ProSciTech, Thuringowa, QLD, Australia) in Et$_2$O in line with the following procedure: freeze-substitution, 24 hr at -100°C, 168 hr at -90°C, 350 hr at -70°C, and 24 hr at -50°C; resin infiltration at -50°C, 1 hr 1:1, 1 hr 1:2, 1 hr 100% HM20, 24 hr 100% HM20, followed by two 48 hr UV polymerization steps at -50 °C and 20°C. The presence of water was minimized by preparing all solutions and performing freeze-substitution with molecular sieve (3A, Acros Organics).

Optic nerve segments were dry serially sectioned with a diamond knife (Histo, Diatome): three 1 μm sections were collected for toluidine blue staining for identification of the injury site; the next ten 1 μm sections were collected in pairs - one for NanoSIMS and the next for immunohistochemical analyses; the following 10 μm were discarded. The pattern was repeated until the whole nerve segment had been serially sectioned. For each optic nerve segment, a single section at the injury site was selected for NanoSIMS analysis, transferred onto 1 inch diameter polished silicon wafers, and flattened/attached using a single drop of dry tetrahydrofuran. The wafers were then coated with 10 nm Au to provide conductivity at high voltage in the NanoSIMS. Adjacent sections for immunohistochemical analyses were stored at room temperature.

**Measurement of samples using NanoSIMS**

For each optic nerve segment, sections were viewed to locate disturbance of the tissue indicating the dorsal injury site and a single section clearly at the injury site was chosen for analysis. One or more fields of view (30 μm x 30 μm) in the central region of the selected section (and therefore in an area vulnerable to secondary degeneration) were analyzed. Fields of view, including both axons and glia, were chosen using light-microscopy images of adjacent Toluidine blue-stained sections, in order to ensure a range of cellular features could be analyzed. Elemental images were acquired using the CAMECA NanoSIMS 50 ion microprobe at The University of Western Australia. Prior to imaging, all fields of view were implanted with a primary ion dose of 1 x 10$^{17}$ ions/cm$^2$, thereby affording a steady state of secondary ion yield. A Cs$^+$ primary ion beam (nominal beam diameter = 100 nm, current = 1.5 pA) was used to sputter the negative ion species $^{12}$C$^{12}$C, $^{12}$C$^{14}$N, $^{31}$P, $^{32}$S, $^{35}$Cl and...
secondary electrons. An O₃ primary ion beam (nominal beam diameter = 600 nm, current = 28 pA) was then used to sputter the positive ion species ¹²C, ²³Na, ³⁹K and ⁴⁰Ca. All fields of view were imaged at a resolution of 256 × 256 pixels, with a dwell time of 30 ms/pixel. There were no significant isobaric interferences, so it was not necessary to tune the mass spectrometer to high mass resolution; charge build-up during Cs⁺ primary beam use was not observed. The combination of underlying Si substrate and overlying Au coat provided adequate sample conductivity. As the valency of the detected ions is not discernible, detected ions are referred to by their isotopic species (e.g. ⁴⁰Ca) and elemental symbol (Ca), not oxidation state (e.g. Ca²⁺).

Analysis of NanoSIMS data

The NanoSIMS produces secondary ion maps with 32-bit data density, where each pixel records the number of secondary ion counts within the acquisition time. Secondary ion images were corrected for detector deadtime (44 ns) and analyzed using the OpenMIMS plugin for Fiji/ImageJ (version 2.0; NIH). For quantitative analysis of ⁴⁰Ca, area (pixels) and mean secondary ion count (counts/pixel) data for specified regions-of-interest within the image were extracted. For analysis, area and mean counts were converted to μm² (72,817 pixels²/μm²).

In order to correct for any signal strength variability from sample to sample, each ion-signal from which data were extracted and analyzed quantitatively was displayed as a ratio of the ¹²C signal collected for the entire field of view.

To allow for easier visualization of structural information, overlays of three secondary ion images, to which each had been assigned one RGB color channel, were constructed (Fig. 1A and B). Ten secondary ion channels combined into 3 channel RGB color images result in 720 unique permutations, shown are two possibilities. Changes in the displayed color were indicative of changes in the ratio of one element relative to others. Overlays of ³¹P, ³₂S and ¹²C¹⁴N secondary ion images (P-S-CN assigned to RGB color channels in order listed) were found to best visually enhance structural information (nuclei, axons, myelin) present in individual images.

Ca microdomains were defined as any area or group of pixels with a ⁴⁰Ca signal in excess of 20 secondary ion counts/pixel, with regions of punctate or ‘speckled’ Ca defined as 0 ≤ 20 secondary ion counts/pixel. Thresholds set to these values were used to generate regions-of-interest using Fiji/ImageJ, allowing data extraction only within, or outside, Ca microdomains.

Microdomain/punctate regions-of-interest were exported as 45 counts within the image were extracted. For analysis, area and mean values were expressed as a proportion of the total area of the tissue type (D, glial = black; axonal = white) can be used to generate 8-bit greyscale maps (E) used to define individual regions of interest in Fiji/ImageJ: example shown (F; non-P co-localized glial microdomains, color code = 666666; green and blue are excluded regions). All images 30 μm × 30 μm, scale bar = 10 μm.

The number of Ca microdomains was counted manually and expressed as a density, relative to the area (μm²) of axonal or glial regions. Clustered Ca microdomains were counted as separate if there was a clearly defined area of lower signal between two points of higher signal. In order to compare the relative areas of non-P or P co-localized Ca microdomains, Ca microdomain areas were expressed as a proportion of the total area of the tissue type (axonal or glial) in which they were found. The total area (of the pre-determined region-of-interest) and the sum of ⁴⁰Ca and of ¹²C signal/pixel (counts/pixel within area of predetermined region-of-interest) were determined using the OpenMIMS plugin. The mean ± S.E.M. size of Ca microdomains were expressed in μm².

Journal Name, [year], [vol], 00–00 | 3
and were calculated by dividing the total area of P or non-P co-localized Ca microdomains by the number of these Ca microdomains. The relative Ca concentration of the Ca microdomains is expressed as mean $^{40}\text{Ca}/^{12}\text{C}$ ratio ± S.E.M/ unit area ($\mu\text{m}^2$) of the tissue type in which the Ca microdomains were found.

Quantification of punctate Ca signal that was not part of Ca microdomains was conducted on whole images. Nuclei were identified with reference to Hoechst stained adjacent sections and P-S-CN overlays, and were excluded from these analyses. The mean $^{40}\text{Ca}/^{12}\text{C}$ ratio ± S.E.M was normalized for the punctate area ($\mu\text{m}^2$) in each image. Visualization of the punctate Ca signal was achieved by adjusting the displayed minimum-maximum grayscale values to correspond to 0 - 60 in 32-bit $^{40}\text{Ca}/^{12}\text{C}$ ratio images (scale factor of 10) generated using the OpenMIMS plugin.

No marked differences in mean densities, size or intensities of Ca microdomains were observed between multiple fields of view within a single optic nerve section, although the nature of the single observations generated per region-of-interest precluded statistical confirmation of their similarity. Therefore, mean values for all outcomes were generated for the 1 - 3 fields of view analyzed/optic nerve (therefore generating one value/animal for each outcome measure, n = number of animals) and grand means and standard errors of the mean (S.E.M.) for all animals/time point were calculated. Of necessity, a microanalysis technique such as NanoSIMS must assess a small fraction of the available tissue section. As the frequency of Ca microdomains is relatively low, there is some variability between replicates. Variation could only have been reduced by substantially increasing the number of fields of view sampled, resulting in a reduction in the number of time points post injury that could be assessed, due to the practical space, time and cost limitations of the analysis technique. The decision was made to maintain the chosen study design and appropriate statistical analyses has enabled interpretation of the data. Due to the relatively small sample size, data was assumed to be normally distributed. Values were statistically analyzed using 2-way ANOVA, ANOVA and Bonferroni/Dunn post-hoc tests (IBM SPSS Statistics Version 20; IBM Corporation, Armonk, NY) as applicable, requiring a significance of $P \leq 0.05$. Lavene’s test of equal variance was used to show that population variances at different time points were not significantly different ($P > 0.05$).

**Immunohistochemistry**

Sections adjacent to NanoSIMS sections were analyzed immunohistochemically according to previously described procedures, with the following modifications. Sections were etched in saturated NaOH/100% EtOH for 2 minutes and graded through an EtOH series followed by microwave antigen retrieval in 10 mM citric acid buffer, pH 6.0. Primary antibodies (diluted in PBS + 0.2% Triton X100 (PBST) containing 5% normal donkey serum) were used to detect: neurons (anti-β-III tubulin; 1:500, Tuj1 clone Covance or ab18207 Abcam); myelin basic protein (anti-MBP; 1:500, ab40390 Abcam); astrocytes (anti-GFAP; 1:500, GA5 Millipore); Ca$^{2+}$/calmodulin-dependent protein kinase II (anti-CaMKII; 1:100, ab2724 Abcam) or voltage-dependent anion channel 2 (anti-VDAC2; 1:200, HPA043475 Sigma). Antibody binding was visualized following 2 h incubation at room temperature with appropriate secondary antibodies (Alexa Fluor 488, 555 or 647 (Molecular Probes) all at 1:400 in PBST). Nuclei were identified using Hoechst stain (10 µg/mL, Invitrogen). Slides were coverslipped using Fluoromount-G (Southern Biotechnology) and visualized and imaged using a Nikon Eclipse Ti inverted microscope (Nikon Corporation, Japan) with a 40X/1.3 N.A. oil immersion objective or a Leica TCS SP2 multiphoton confocal microscope (Leica Microsystems Heidelberg GmbH, Germany) with a PL APO CS 63.0X/1.40 oil immersion objective. All images collected using Nikon Elements AT software were deconvoluted using autoquant blind deconvolution. All sections for each antibody were immunostained at the same time to ensure uniformity of the immunohistochemical procedures, and images were captured at constant exposure times. Control sections stained only with secondary antibodies were included in all experiments and showed no or only minimal fluorescence (data not shown).

**Results**

NanoSIMS characterization of Ca microdomains in normal optic nerve

In normal optic nerve excised from uninjured animals, NanoSIMS secondary ion imaging reveals the distribution of Ca either as localized, high intensity circular features as small as 1 μm in diameter, henceforth referred to as Ca microdomains (Fig. 2A) or, as a disseminated, low intensity, background signal with a speckled appearance, referred to as punctate. A signal threshold was used to delineate Ca microdomains for...
in the various forms of Ca microdomains is illustrated (Fig. 3).

Structural information can be visualized in images generated from secondary electrons and from RGB overlays of $^{31}$P, $^{32}$S and $^{12}$C$^{14}$N (P-S-CN) signal in the same field of view (C, D). P-S-CN images also allow for identification of areas enriched in P (E, pink dot), which in some cases co-localize with Ca microdomains (F, overlaid in G), circled in yellow. Light microscopy (H) along with immunohistochemical staining (I) of sections encompassing the NanoSIMS field of view (boxed in yellow) allow visualization of structural features that were identified as glial regions containing GFAP$^+$ astrocytes (green) or axonal regions containing β-III tubulin$^+$ (red) axonal profiles associated with MBP$^+$ myelin (blue) (I). Gliarial (hashed in yellow) and axonal regions can then be identified in P-S-CN images (D). Images of Ca microdomains overlaid on P-S-CN images (I) together with immunohistochemical staining for VDAC2 (K), demonstrate co-localization of some Ca microdomains (green) with VDAC2 (purple) (L). Both non-P co-localized (circled in yellow) and P co-localized (circled in red) Ca microdomains were found to associate with VDAC2. Images of Ca microdomains overlaid on P-S-CN images (M) together with immunohistochemical staining for CaMKII (N) show that Ca microdomains (green, non-P co-localized circled in yellow, P co-localized circled in red) do not appear to co-localize with CaMKII (purple) (O). Images H and I, scale bar = 30 μm. All other NanoSIMS images 30 μm × 30 μm, scale bar = 10 μm.

quantification purposes (Fig. 2B). Secondary electron NanoSIMS images (Fig. 2C) allowed visualization of morphological regions that corresponded to specific colored regions in RGB overlays of $^{31}$P, $^{32}$S and $^{12}$C$^{14}$N (P-S-CN) images (Fig. 2D), indicating differences in elemental composition. Some Ca microdomains were co-localized with areas of enriched $^{31}$P signal, observed as small pink dots in P-S-CN images (Fig. 2E, co-localization illustrated in Fig. 2F, G, larger microdomain of those circled in yellow), whereas others did not (Fig. 2E - G, smaller microdomain of those circled in yellow). Ca microdomains that were or were not co-localized with P-enriched areas (referred to henceforth as P or non-P co-localized) were quantified separately for axonal and glial regions (see below). Distribution of calcium in the various forms of Ca microdomains is illustrated (Fig. 3).

![Fig. 3 Schematic illustrating the distribution of Ca in P- and non-P co-localized microdomains in axonal and glial regions.](image)

We first investigated the sub-cellular localization of P and non-P Ca microdomains within the normal optic nerve. Ca$^{2+}$ microdomains have been proposed to be associated with the plasma membrane $^{11}$ and intracellular structures including endoplasmic reticulum and mitochondria $^{13}$. To confirm this using our technique and to additionally determine whether P and non-P co-localized Ca microdomains are located in different sub-cellular regions, P-S-CN images overlaid with $^{40}$Ca/$^{12}$C images (Fig. 2J, M) were compared to adjacent sections immunohistochemically identifying VDAC2 (labels the mitochondrial membrane; Fig. 2K, L) or CaMKII (located at the plasma membrane when activated, Fig. 2N, O). As a kinase, we hypothesized that CaMKII may be selectively associated with P co-localized Ca microdomains. Immunohistochemical analyses could not be conducted on the same sections imaged using NanoSIMS due to the required Au coating of the sample. However, comparison of adjacent sections allows an indication of possible co-localization of VDAC2 and/ or CaMKII with the Ca microdomains. The majority of Ca microdomains in normal optic nerve were closely associated but not precisely co-localized with VDAC2 (circled in yellow for non-P and in red for P co-localized areas), consistent with Ca accumulation in mitochondria $^{12}$; there was no indication of selective association for non-P or P co-localized Ca microdomains. Quantitative analysis of co-localization between NanoSIMS Ca microdomains and VDAC2 staining would require confocal imaging and analysis of a single image in the z plane. As such, it is not possible to conduct co-localization analyses in the current study as immunohistochemical and NanoSIMS analyses can only be conducted on adjacent sections. Furthermore, we did not observe precise co-localization, so co-localization values are likely to be negligible. There appeared to be no consistent association of either non-P co-localized (circled yellow) or P co-localized (circled red) Ca microdomains with CaMKII.

Light microscopy images and immunohistochemical analyses of sections adjacent to those used for NanoSIMS analysis also allowed for differentiation of glial and neuronal cell types (Fig. 2H, I). The relatively featureless areas in the secondary electron images and darker areas in the P-S-CN images corresponded to areas containing GFAP$^+$ astrocytes, a subset of glial cells (Fig. 2D, hashed in yellow, determined with reference to Fig. 2H and I), whereas circular features correspond to β-III tubulin$^+$ axons closely associated with MBP$^+$ myelin (Fig. 2D, I). Quantification and measurement of non-P and P co-localized Ca microdomains was therefore conducted separately in axonal and glial regions, based on comparisons with P-S-CN images of the same field of view (Fig. 4).

To compare the proportion of non-P and P co-localized Ca microdomains within each tissue type (glial and axonal), the area of Ca microdomains was expressed as a proportion of the total area of the tissue type in which the Ca microdomains were found (glial or axonal). In normal optic nerve, glial regions contained a significantly greater proportion of non-P co-localized Ca microdomains than P-localized Ca microdomains (Fig. 4A, P ≤ 0.05), while there was no significant difference between the microdomains in axonal regions (Fig. 4B, P > 0.05). Comparison of the proportion of Ca microdomains between the tissue types revealed that there was a significantly greater proportion of non-P co-localized Ca microdomains in glial regions than axonal regions (Fig. 5A, P ≤ 0.05).

Density of a subtype of Ca microdomains was reduced during secondary degeneration

To directly quantify changes in Ca microdomains vulnerable to spreading damage following neurotrauma, we assessed the density of Ca microdomains in glial and axonal regions from central optic nerve, undergoing secondary degeneration following dorsal partial nerve injury (Fig. 4C-H) $^{30, 31}$. In glial regions, the total density of Ca microdomains (both non-P and P co-localized) significantly decreased at 5 minutes, 1, 3, 7 days and 1 month.
Fig. 4 Histograms show mean Ca microdomain proportions ± S.E.M. in glial (A) and axonal (B) regions. Representative NanoSIMS images illustrate the changes in various microdomain types observed in normal optic nerve (C) and at 1 day following partial optic nerve transection (D). Scale bar = 10 μm. Dot plots show quantification of mean non-P (E) and P (F) co-localized Ca microdomain density in glial regions, and non-P (G) and P (H) co-localized Ca microdomain density in axonal regions (number/μm²). Significant differences compared with normal optic nerve are indicated (* P ≤ 0.05). Black circles indicate mean density for each animal at a particular time point, grey vertical bars illustrate the range and red horizontal bars indicate grand means for 5 each time point. Histograms show: size of non-P and P co-localized Ca microdomains in glial (I) and axonal (J) regions (in μm²). The mean 40Ca/12C ratio ± S.E.M. of non-P and P co-localized Ca microdomains in glial (K) and axonal (L) regions (in arbitrary units). Significant differences between values at each time-point are indicated by # (P ≤ 0.05) and significant changes as a consequence of injury are indicated by * (P ≤ 0.05).

after injury, compared to normal optic nerves (control from uninjured normal animals) (normal 0.058 ± 0.017; 5 minutes 0.016 ± 0.006; 1 day 0.013 ± 0.005; 3 day 0.032 ± 0.009; 7 day 0.018 ± 0.009; 1 month 0.014 ± 0.005, P ≤ 0.05; 3 month 0.044 ± 0.012, P > 0.05). When the densities of non-P co-localized Ca microdomains were analysed separately, significant decreases were observed at 5 minutes, 1, 3, 7 days, 1 and 3 months after injury, compared to normal optic nerves (Fig. 4E, P ≤ 0.05). No significant differences in glial P co-localized Ca microdomains were observed at any of the time points assessed after injury (Fig. 4F, P > 0.05). In axonal regions, a similar trend was observed: the density of non-P co-localized Ca microdomains significantly decreased at 5 minutes, 1 and 7 days post injury (Fig. 4G, P ≤ 0.05) and there were no significant changes in the density of P co-localized axonal microdomains compared to normal (uninjured control) optic nerves (Fig. 4H, P > 0.05). Total Ca microdomain density in axonal regions (sum of non-P and P co-localized) did not change following injury (data not shown, P > 0.05); indicating that pooling the P and non-P co-localized Ca microdomains together masked the significant changes observed in the non-P co-localized microdomains in axonal regions. The described changes in Ca microdomain density can be observed in representative images in Figure 4C-D, with noticeably fewer microdomains at 1 day after injury (Fig. 4D) compared to normal optic nerve (Fig. 4C).

Changes in the proportion of non-P and P co-localized Ca microdomains following partial optic nerve transection were consistent with the changes in density described above. Post injury, the glial regions exhibited a significant decrease in proportion of non-P Ca microdomains at all time points up to 3 months (Fig. 4A, P > 0.05), while the P co-localized microdomains remained at normal levels at all time points after injury. As such, the proportion of non-P co-localized Ca microdomains was no longer greater than the proportion of P co-localized microdomains until 3 months after injury. Note that the apparently greater proportion of non-P than P co-localized Ca microdomains in glial regions at day 3 (Fig. 4A) did not reach statistical significance due to the variability inherent in the data.
set as a whole. The proportion of axonal non-P or P co-localized Ca microdomains did not change significantly, relative to normal optic nerve, at any time point post injury (Fig. 4B, P > 0.05).

As a consequence of the reduction in the proportion of non-P co-localized Ca microdomains in glia after injury, there was no longer a significant difference in the proportions of these microdomains between axons and glia (Fig. 5A, P > 0.05). The only significant difference between glial and axonal regions in terms of the proportion of P co-localized Ca microdomains, was a slightly greater proportion in axonal Ca microdomains at day 3 (Fig. 5B, P ≤ 0.05). There were no apparent differences in the association of P or non-P lo-localized Ca microdomains with VDAC2 or CaMKII after injury (Fig. 6, day 7 shown compared to Fig. 2J-O).

Assessment of the size of non-P and P co-localized Ca microdomains assessed individually for glial and axonal compartments (Fig. 4I, J) showed that there were no significant changes in the mean size of Ca microdomains as a result of partial optic nerve transection (Fig. 4I, J, P > 0.05), illustrated in representative images at 1 day after injury (Fig. 4D) compared to normal optic nerve (Fig. 4C). There were also no significant differences between the size of non-P compared to P co-localized Ca microdomains in normal optic nerve or at any time after injury (Fig. 4I, J, P > 0.05), or between the size of Ca microdomains in axonal compared to glial regions (Fig. 5C, D, P > 0.05). Note that the size of non-P co-localized Ca microdomains in glial regions at day 1 after injury was reduced when ANOVA was confined to non-P co-localized microdomains (Fig. 5C, P ≤ 0.05). It is worth noting that the O beam used to detect Ca in the NanoSIMS is larger than the Cs beam used to detect the P-S-CN structural information. Therefore, the Ca microdomains may appear larger in our NanoSIMS images than they actually are.

The mean $^{40}\text{Ca}/^{12}\text{C}$ ratio was assessed as an indication of the amount of Ca in the microdomains. A significant increase was observed in the Ca in axonal non-P co-localized Ca microdomains at 7 days post injury, relative to normal optic nerve (Fig. 4L, P ≤ 0.05); no differences were observed in glial non-P co-localized Ca with injury (Fig. 4K, P > 0.05). No significant changes in Ca were observed for glial or axonal P co-localized Ca microdomains following injury (Fig. 4K, L, P > 0.05). Ca in P co-localized Ca microdomains was not different to that in non-P co-localized in normal optic nerve or at any time after injury for

---

**Fig. 5** Histograms show: mean Ca microdomain proportions ± S.E.M. in non-P (A) and P (B) co-localized Ca microdomains; size of glial and axonal non-P (C) and P co-localized (D) Ca microdomains (in μm²). The mean $^{40}\text{Ca}/^{12}\text{C}$ ratio ± S.E.M. of glial and axonal non-P (E) and P co-localized (F) Ca microdomains (in arbitrary units). Significant differences between values at each time-point are indicated by # (P ≤ 0.05) and significant changes as a consequence of injury are indicated by * (P ≤ 0.05).

**Fig. 6** Images of Ca microdomains overlaid on P-S-CN images (A) together with immunohistochemical staining for VDAC2 (B), indicate co-localization of some Ca microdomains (green) with VDAC2 (purple) (C). Both non-P co-localized (circled in yellow) and P co-localized (circled in red) Ca microdomains were found to associate with VDAC2. Images of Ca microdomains overlaid on P-S-CN images (D) together with immunohistochemical staining for CaMKII (E) show that Ca microdomains (green, non-P co-localized circled in yellow, P co-localized circled in red) do not appear to co-localize with CaMKII (purple) (F), scale bar = 10 μm.
either axonal or glial regions (Fig. 4K, L, P > 0.05). Similarly, Ca in axonal regions was not different to that in glial regions for either non-P or P co-localized Ca microdomains (Fig. 5E, F, P > 0.05). The mean punctate \(^{40}\text{Ca}/^{12}\text{C}\) ratio was quantified in normal optic nerve and at each time point post injury, but no significant differences were observed (P >0.05, data not shown).

**Discussion**

Ca\(^{2+}\) changes as a consequence of injury to the CNS are complex, rapid and result in cellular degeneration and death, but have generally been viewed at a whole cell rather than at a Ca\(^{2+}\) microdomain level. Using NanoSIMS combined with immunohistochemical identification of individual cell types in *ex vivo* tissue samples, we have identified novel sub-populations of Ca microdomains that are or are not co-localized with areas enriched in P. We show for the first time that non-P and P co-localized microdomains into the intra- and/or optic nerve and at each time point post injury, but no significant

**Fig. 7** Model illustrating potential consequences of Ca microdomain changes in axonal and glial regions early post injury (1 day), compared to normal optic nerve. There is a decrease in the number of both axonal and glial non-P co-localized microdomains with injury, which may result in increased release of free (punctate) Ca into the cytosol and/ or extracellular space.

The combination of NanoSIMS with immunohistochemistry or cell labeling techniques has recently been reported, but to our knowledge, ours is the first study combining direct high resolution NanoSIMS visualization of Ca microdomains with immunohistochemical identification of individual cell types in *ex vivo* mammalian tissue samples. This new technique allows for the measurement and quantification of Ca\(^{2+}\) microdomains in specific cell types in tissue and allowed demonstration that the changes in Ca microdomains were more pronounced in glial than axonal compartments, perhaps reflecting the reactivity of astrocytes following injury to the CNS and propagation of Ca\(^{2+}\) waves through the astrocytic syncretum. Ca\(^{2+}\) microdomains can be transient phenomena, however the rapid cryopreservation of tissue for NanoSIMS analysis allows their accurate visualization at a particular point in time. Nevertheless, a limitation of our technique is that real-time assessments of Ca\(^{2+}\) signaling dynamics cannot be conducted. We also cannot guarantee that the sample preservation has not affected the chemistry of the samples, but we are confident in the results given our rapid cryopreservation technique, combined with low-temperature substitution and embedding.

In *vitro* investigations designed to model the injury scenario have revealed that release of excess, localized Ca\(^{2+}\) from Ca\(^{2+}\) microdomains, together with Ca\(^{2+}\) influx from the extracellular space, leads to increased intracellular Ca\(^{2+}\) concentrations, oxidative stress and death of neurons and glia. The role played by Ca\(^{2+}\) microdomains in the context of secondary injury in the CNS has not been studied extensively, perhaps due to the difficulty in studying Ca\(^{2+}\) microdomains *in situ* as opposed to global Ca\(^{2+}\) levels. The decrease in Ca microdomain density we have observed in axonal and glial regions vulnerable to secondary degeneration would suggest a decrease in overall Ca\(^{2+}\) sequestered in microdomains, with subtle differences between cell types.

Given the overall increase in Ca\(^{2+}\) concentration that has been reported in injured spinal cord, particularly at the impact site, our data suggest an efflux of Ca\(^{2+}\) from microdomains into a global punctate distribution, similar to that which has been observed from Ca\(^{2+}\) microdomains associated with mitochondria, sarcoplasmic and endoplasmic reticulum in cultured HeLa cells and myocytes. Released Ca\(^{2+}\) could exist intra- or extracellularly and the current NanoSIMS analyses do not distinguish between these; increasing the lateral resolution may increase sensitivity for this outcome in future assessments. The NanoSIMS acquisition parameters for the current study were designed to allow quantification of Ca microdomains rather than the punctate \(^{40}\text{Ca}/^{12}\text{C}\) signal; adjustments to sensitivity may reveal alterations in punctate \(^{40}\text{Ca}/^{12}\text{C}\) ratio with injury in future studies. As such, here, we were not able to quantitatively detect increases in punctate \(^{40}\text{Ca}/^{12}\text{C}\) signal in optic nerve vulnerable to secondary degeneration, consistent with release of Ca\(^{2+}\) from microdomains. It is important to note that there may have been an initial increase in Ca microdomain density in nerve vulnerable to secondary degeneration immediately following injury, which declined to lower levels than normal by 5 minutes after injury. Further, it is possible that released Ca\(^{2+}\) may focus at the injury site, away from the area of secondary degeneration analyzed in the current study. However, given that we have previously...
reported no difference in mean Ca signal between areas affected by the primary injury and those vulnerable to secondary degeneration. It is possible that the total Ca²⁺ concentration actually falls throughout optic nerve following partial injury, implying that the reported decrease in extracellular Ca²⁺ following spinal cord injury is not compensated for by intracellular Ca²⁺ concentration increases, as has been previously suggested. Future studies will investigate these possibilities.

Conclusions
NanoSIMS analyses of Ca microdomains have provided novel insights regarding association with P and allowed quantitative, direct measures of complex Ca²⁺ microdomain changes in glial and axonal regions vulnerable to secondary degeneration following neurotrauma. The study has also illustrated the utility of this technology, applicable to a wide range of mammalian systems in both normal and injured states.

Notes and references

*BioNano, School of Chemistry and Biochemistry, The University of Western Australia, 35 Stirling Hwy, Crawley WA 6009, Australia
*Experimental and Regenerative Neurosciences, School of Animal Biology, The University of Western Australia, 35 Stirling Hwy, Crawley WA 6009, Australia
Centre for Microscopy, Characterisation and Analysis, The University of Western Australia, 35 Stirling Hwy, Crawley WA 6009, Australia

*To whom correspondence should be addressed: Melinda Fitzgerald, Experimental and Regenerative Neurosciences, School of Animal Biology M317, The University of Western Australia, 35 Stirling Hwy, Crawley WA 6009, Australia. Tel: 61 8 6488 2353, Fax: 61 8 6488 7527, Email: lindy.fitzgerald@uwa.edu.au

33. O. Hommou and W. Young, in The axon: structure function and pathophysiology, eds. S. G. Waxman, J. D. Kocsis and P. K. Stys, this journal is © The Royal Society of Chemistry [year].
Acknowledgements

This work was supported by the Neurotrauma Research Program (Western Australia) and NH&MRC (Grant ID: APP1028681). Prof. Dunlop is a Principal Research Fellow of the NH&MRC, Australia (Grant ID: APP1002347). We thank Associate Professor Nathan J. Pavlos, The University of Western Australia, for kindly providing us with a sample of VDAC2 antibody; Mr Guy Ben-Ary of Cell Central, The University of Western Australia, for assistance with fluorescent microscopy; Mr John Murphy, The University of Western Australia, for assistance with confocal microscopy and Associate Professor Jenny Rodger for critical review of the manuscript. The authors acknowledge the facilities, scientific and technical assistance of the Australian Microscopy and Microanalysis Research Facility at the Centre for Microscopy, Characterisation and Analysis, The University of Western Australia, a facility funded by the University, State and Commonwealth Governments. This work was made possible in part by the OpenMIMS software whose development is funded by the NIH/NIBIB National Resource for Imaging Mass Spectrometry, NIH/NIBIB 5P41 EB001974-10.