1	Comparison of ion channel inhibitor combinations for limiting secondary degeneration
2	following partial optic nerve transection
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23 Abstract

Following neurotrauma, secondary degeneration of neurons and glia adjacent to the injury 24 leads to further functional loss. A combination of ion channel inhibitors (lomerizine + oxATP 25 + YM872) has been shown to be effective at limiting structural and functional loss due to 26 secondary degeneration. Here we assess efficacy of the combination where oxATP is 27 replaced with Brilliant Blue G (BBG), a more clinically applicable P2X₇ receptor inhibitor. 28 Partial optic nerve transection was used to model secondary degeneration in adult female rats. 29 Animals were treated with combinations of lomerizine + YM872 + oxATP or lomerizine + 30 31 YM872 + BBG, delivered via osmotic mini pump directly to the injury site. Outcomes assessed were Iba1+ and ED1+ microglia and macrophages, oligodendroglial cell numbers, 32 node/paranode structure and visual function using the optokinetic nystagmus test. The 33 34 lomerizine + BBG + YM872 combination was at least as effective at the tested concentrations as the lomerizine + oxATP + YM872 combination at preserving 35 node/paranode structure and visual function when delivered locally. However, neither ion 36 channel inhibitor combination significantly improved microglial/macrophage nor 37 oligodendroglial numbers compared to vehicle treated controls. In conclusion, a locally 38 39 delivered combination of ion channel inhibitors incorporating lomerizine + BBG + YM872 is 40 at least as effective at limiting secondary degeneration following partial injury to the optic 41 nerve as the combination incorporating oxATP.

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Keywords: secondary degeneration; neurotrauma; ion channel inhibitor; myelin; visual
function

47 Introduction

Following neurotrauma, a series of metabolic and structural changes are propagated in 48 initially undamaged tissue, associated with increased intracellular Ca²⁺, oxidative stress and 49 50 apoptotic cell death of neurons and glia (Dong et al. 2009). Since the initial insult is often unavoidable, treatments for functional recovery after neurotrauma focus heavily on limiting 51 this secondary damage (Doan et al. 2016). However, despite extensive research, effective 52 pharmacotherapeutic treatments for secondary degeneration following neurotrauma are 53 limited (Kwon et al. 2011). In order to successfully limit secondary degeneration following 54 55 neurotrauma, it is important to test efficacy of treatments in appropriate animal models of injury. Partial optic nerve transection is an established and useful model for investigating 56 secondary degeneration, where the dorsal optic nerve of adult rats is partially transected, 57 58 allowing for spatial separation between the primary and subsequent secondary degeneration (Levkovitch-Verbin et al. 2003; Blair et al. 2005). The model has been further characterised 59 and employed to assess efficacy of pharmacotherapeutics for secondary degeneration, 60 delivered directly to the injury site using osmotic mini-pumps (Fitzgerald et al. 2009a; 61 Fitzgerald et al. 2009b; Savigni et al. 2013; O'Hare Doig et al. 2017). 62 63 Secondary degeneration is characterised by a myriad of reactive metabolic pathways, including inflammation, excitotoxicity, mitochondrial dysfunction and oxidative stress, 64

associated with structural deficits, dysmyelination and apoptotic cell death (Tymianski and

66 Charles 1996; Dong et al. 2009; Maxwell 2013). Ca^{2+} overload is considered to be a major

67 trigger for the toxic mechanisms of secondary degeneration (Farooqui et al. 2008). Using the

68 partial optic nerve transection model, we have previously demonstrated that a locally

69 delivered combinatorial treatment strategy to limit excess Ca^{2+} influx through voltage gated

calcium channels, P2X₇ receptors and Ca^{2+} permeable AMPA receptors with lomerizine,

oxATP and YM872 respectively, reduced myelin decompaction, preserved node/paranode

structure and visual function (Savigni et al. 2013). Acute outcomes indicated that early
preservation of node/paranode structure and OPC numbers was associated with longer term
preservation of visual function (O'Hare Doig et al. 2017).

75 Following mild traumatic brain injury, it is currently unclear as to whether there is a blood brain barrier breach, with studies reporting varying degrees of compromise of blood brain 76 barrier integrity following injury (Deford et al. 2002; Tomkins et al. 2011; Zetterberg et al. 77 2013). Therefore, pharmacotherapies designed to treat all but the most severe cases of 78 neurotrauma, need to be able to travel across the closed blood brain barrier following 79 80 systemic delivery. oxATP does not appear to be able to cross the blood brain barrier (Peng et al. 2009). Thus, for clinical applicability of the lomerizine, oxATP and YM872 combination, 81 82 oxATP needs to be substituted with a blood brain barrier permeable P2X7 receptor 83 antagonist. Here we introduce an alternative P2X₇ receptor inhibitor Brilliant Blue G (BBG) to the combination. BBG has previously shown therapeutic effects following neurotrauma 84 (Peng et al. 2009; Kimbler et al. 2012; Wang et al. 2015), and importantly, can cross the 85 closed blood brain barrier (Wong et al. 2011). This study compared the efficacy of the ion 86 channel inhibitor combination of lomerizine + BBG + YM872 to a combination with 87 88 lomerizine + oxATP + YM872 for limiting secondary degeneration and restoring function 89 following partial optic nerve transection.

90

91 Methods

92 Animals and study design

93 Thirty-eight adult, female PVG rats were obtained from the Animal Resource Centre in
94 Murdoch, Western Australia. The animals were housed under 12-hour light/dark cycles with
95 ad libitum access to food and water. All procedures were approved by the University of

Western Australia Animal Ethics Committee (approval number RA3/100/1485) and were in accordance with the National Health and Medical Research Council (NHMRC) of Australia Code of Practice for use of Animals for Scientific Purposes. The animals were divided into four experimental groups, a sham injured, vehicle treated group (n = 8); an injured, vehicle treated group (n = 10); an injured, lomerizine + oxATP + YM872 treated group (n = 10); and an injured, lomerizine + BBG + YM872 treated group (n = 10); with the sham group serving as an uninjured, vehicle treated control.

103 Surgical procedures

Partial optic nerve transection (day 1) and left eyelid suturing (day 3) were performed as 104 previously described (Fitzgerald et al. 2009a), under Ketamine (Ketamil, 50mg/kg, Troy 105 Laboratories) and Xylazine (Ilium Xylazil, 10mg/kg, Troy Laboratories) anaesthesia 106 107 administered intraperitoneally. In brief, for the partial transection surgery: the skin overlying the skull behind the right eye was incised. The optic nerve was accessed and the nerve 108 parenchyma exposed by making a longitudinal cut in the sheath using fine iridectomy 109 scissors. About 1mm behind the right eye, the dorsal aspect of the optic nerve was partially 110 lesioned to a depth of approximately 200µm with a diamond radial keratotomy knife 111 112 (Geuder); the depth determined by the protrusion of the blade beyond the surrounding guard. Sham injury included all procedures except the cut in the sheath and the partial optic nerve 113 lesion. Surgical implantation of Alzet osmotic mini-pumps was performed as described 114 (Savigni et al. 2013). Immediately following surgery, subcutaneous injections of analgesia 115 (2.8mg/kg carprofen, Norbrook) and 1mL sterile phosphate buffered saline (PBS) were 116 administered. 117

118 Treatments

Lomerizine (30mg/kg, LKT Labs©) was orally administered in butter vehicle twice daily 8 119 hours apart, until end of experiment, commencing once animals were ambulatory following 120 surgery as previously described (Fitzgerald et al. 2009a). oxATP (1mM), BBG (540µM) and 121 122 YM872 (240µM) were delivered via osmotic mini-pump at 0.5µL/h in PBS vehicle. Concentrations employed for oxATP and YM872 were consistent with our previous studies 123 where efficacy was demonstrated (Savigni et al. 2013), and the BBG dose was chosen with 124 reference to the literature describing efficacy of BBG and YM872 in related models 125 (Takahashi et al. 2002; Diaz-Hernandez et al. 2012; Cervetto et al. 2013). The sham injured 126 127 and partial optic nerve transection injured, vehicle treated experimental groups both received PBS via osmotic mini-pump and butter orally, administered as described for the inhibitor 128 treated groups. 129

130 Behavioural assessment

On day 3, the animals were anaesthetised as described above and their uninjured left eyelids 131 132 sutured shut. The optokinetic nystagmus assessment of visual function was performed on day 4 for all animals, in accordance with established procedures (Fitzgerald et al. 2010b). 133 Animals were videoed and number of responses per unit time engaged in the task was 134 135 determined by a single investigator blinded to animal identity. Note that due to a procedural error, n = 5 for the injured, vehicle treated group. Responses were categorised as either 136 smooth pursuits or fast resets. Smooth pursuits are characterised as an elongated head 137 138 rotation tracking the stripes, and fast resets as a rapid, realigning head movement; both elements are an indication of visual ability of the animal (Abdeljalil et al. 2005). 139

140 *Tissue processing and immunohistochemistry*

141 Immediately following behavioural assessment, rats were euthanised with pentobarbitone

sodium (160mg/kg, Delvet), transcardially perfused with 0.9% saline, followed by 4%

143 paraformaldehyde (Sigma-Aldrich) in 0.1M PBS. Optic nerves were dissected and fixation continued overnight by immersion in 4% paraformaldehyde. Tissue was transferred into 15% 144 sucrose (Chem Supply) in PBS, then cryosectioned in longitudinal orientation at a thickness 145 of 14µm and collected onto Superfrost Plus glass microscope slides. Immunohistochemistry 146 was conducted in accordance with established procedures(Fitzgerald et al. 2010a) using 147 primary antibodies recognising: microglial activation markers Iba1 (1:500; Abcam, goat 148 Ab5076) and ED1 (1:500; Merck Millipore, mouse MAB1435); oligodendroglial indicators 149 oligodendrocyte transcription factor 2 (Olig2; 1:500; R&D Systems, goat AF2418) and 150 151 platelet-derived growth factor alpha receptor (PDGFaR; 1:500; Abcam Ab96806); and for paranode and node of Ranvier structures Caspr (1:500; Abcam, rabbit Ab34151), and β -III 152 tubulin (1:500; Merck Millipore, mouse MAB1637). Antibodies were diluted in PBS 153 154 containing 0.2% Triton[™] X-100 and 5% normal donkey serum. Secondary antibodies were Alexa Flour 488 or 555 (1:400; Thermo Fisher Scientific[™]), together with Hoechst 3342 155 (1:1000; Thermo Fisher Scientific[™]) diluted in PBS containing 0.2% Triton[™] X-100. 156 Finally, the sections were mounted and cover slipped using Fluoromount-G (Thermo Fisher 157 Scientific). 158

159 Imaging and Analysis

The ventral optic nerve directly below the site of injury was visualised, with one field of view 160 from one section per animal imaged for each outcome measure. The slides were viewed using 161 either a Nikon Ni-E confocal fluorescence microscope (Nikon Corporation) or a Nikon 162 Eclipse Ti inverted microscope. A series of 13 optical images were taken at 0.5µm 163 increments along the z-axis, and deconvoluted using Nikon Elements AT software. Imaging 164 for each outcome measure was performed in a single sitting with consistent capture settings. 165 All image analysis was performed on Fiji image processing software (NIH) by a single 166 investigator blinded to section identity. Due to poor fixation and tissue processing in a few 167

168	animals, numbers of animals per group analysed for immunohistochemistry outcomes were:
169	sham injured, vehicle treated group $(n = 8)$; injured, vehicle treated group $(n = 10)$; injured,
170	lomerizine + $oxATP$ + YM872 treated group (n = 8); and injured, lomerizine + BBG +
171	YM872 treated group (n =9).
172	Total numbers of Iba1+ resident reactive microglia, ED1+ activated microglia/macrophages,
173	$Olig2+ oligodendroglia and Olig2+/PDGF\alpha R+ oligodendrocyte precursor cells (OPCs) were$
174	counted within a region of interest in a 20x image of the ventral nerve directly beneath the
175	primary injury site and expressed as the mean number of cells/mm ² . For node/paranode
176	analyses, a single 60x z-stack image per animal was divided into a 3 x 3 grid and all
177	complexes with clearly defined Caspr immunostaining in a single randomly selected grid
178	square assessed; at least 30 node/paranode complexes were analysed per animal. Outcome
179	measures were the length of the paranodal gap, defined as the distance between two Caspr+
180	areas; paranode length, as defined by the length of Caspr+ areas; and the percentages of
181	atypical nodal complexes, as previously described (Szymanski et al. 2013).

182 *Statistics*

Results were analysed using IBM SPSS software. Outliers were detected using the Tukey 183 Outlier Detection Model, whereby data that were greater than 1.5 interquartile ranges outside 184 of the first and third quartiles were considered outliers and removed from the data set. No 185 more than two outliers were removed per treatment group for each outcome measure, details 186 provided in figure legends. A single one-way ANOVA encompassing all four treatment 187 groups were performed for each outcome measure. Normality was assumed, Levene's test 188 189 was used to assess homogeneity of variances for each data set ($\alpha = 0.05$). If Levene's test showed equal variance, the Tukey post-hoc was used; for unequal variance, the Games-190 191 Howell post-hoc was applied; both used $p \le 0.05$ to indicate statistical significance.

192

193 **Results**

194 Effects of ion channel inhibitor combinations on microglia and macrophages

195 Numbers of inflammatory cells in ventral optic nerve vulnerable to secondary degeneration

196 were quantified using Iba1 for resident microglia, ED1 for infiltrating

197 microglia/macrophages, and colocalised Iba1+/ED1+ for infiltrating microglial cells (Wu et

al. 2005). The numbers of Iba1+ activated resident microglia were different in the various

treatment groups (Figure 1a; F = 3.483, DF = 3, p = 0.028), with the only statistically

significant difference an increase in the number of Iba1+ cells when animals were treated

with lomerizine + BBG + YM872 compared to the sham injured, vehicle treated group (p =

0.029). There was a trend towards increased Iba1+ cells with injury when comparing the

injured, vehicle treated group with the sham injured, vehicle treated group (p = 0.071). There

204 was no significant difference between the two ion channel inhibitor combinations in the

number of Iba+ cells (p = 0.938).

206 In contrast, there was a significant difference in numbers of ED1+ infiltrating microglia/ 207 macrophages (Figure 1b; F = 2.5, DF = 3, p = 0.079), with the numbers of the injured, vehicle treated group significantly increased compared to sham injured, vehicle treated animals (p =208 0.049). The number of ED1+ cells in the groups treated with either of the ion channel 209 210 inhibitor combinations were not significantly different to either the injured, vehicle treated group or the sham injured, vehicle treated group (p > 0.05). There was no significant 211 difference between the two ion channel inhibitor combinations in the number of ED1+ cells 212 (p = 0.994).213

Similarly, the number of Iba1+/ED1+ infiltrating microglial cells differed with experimental treatment (Figure 1c; F = 2.912, DF = 3, p = 0.041). A significant increase in the numbers of Iba1+/ED1+ cells was observed in the injured, vehicle treated group compared to the sham injured, vehicle treated group (p = 0.024). Neither of the ion channel inhibitor treatment groups had significantly reduced numbers of Iba1+/ED1+ cells compared to the injured, vehicle treated group (p > 0.05). There was no significant difference between the two ion channel inhibitor combinations in the number of Iba1+/ED1+ cells (p = 0.988). Representative images of Iba1+ cells, ED1+ cells and Iba1+/ED1+ cells are shown (Figure 1d).

223 Effects of ion channel inhibitor combinations on oligodendroglia

The densities of Olig2+ oligodendroglia were not different in any of the experimental groups 224 (Figure 2a; F = 2.042, DF = 3, p = 0.128), example cell shown (Figure 2b). However, when 225 the analysis was refined to include immunoreactivity to PDGF α R, thereby detecting OPCs 226 227 more specifically, there were significant differences between experimental groups (Figure 2c, d; F = 4.681, DF = 3, p = 0.008). As expected (O'Hare Doig et al. 2017), there was a 228 significant decrease in the number of OPCs in the injured, vehicle treated group compared to 229 230 the sham injured, vehicle treated group (p = 0.007). However, neither ion channel inhibitor combination groups had significantly increased numbers of OPCs compared to the injured, 231 232 vehicle treated group (p > 0.05), and there was no significant difference between the two ion channel inhibitor combinations in the number of OPCs (p = 0.599). 233

234 Effects of ion channel inhibitor combinations on node/paranode complexes

Significant differences in the length of the paranode between experimental groups were observed (Figure 3a; F = 52.445, DF = 3, p = 0.0001). Partial optic nerve transection resulted in a significant increase in paranode length in the injured, vehicle treated group compared to the sham injured, vehicle treated group (p = 0.0001), as expected from previous studies (Szymanski et al. 2013). While treatment with lomerizine + oxATP + YM872 reduced 240 paranode length (p = 0.0001), paranodes remained longer than in the sham injured, vehicle treated group (p = 0.0001). Treatment with the lomerizine + BBG + YM872 combination also 241 resulted in significantly reduced paranode lengths compared to the injured, vehicle treated 242 group (p = 0.0001), to levels significantly different to the sham injured, vehicle treated group 243 (p = 0.001). Largely similar outcomes were observed when measuring the length of the 244 paranodal gap, indicative of the length of the node of Ranvier (Figure 3b; F = 20.367, DF = 3, 245 246 p = 0.0001). There was no significant difference between the two ion channel inhibitor combinations in the length of the paranode (p = 0.084) or the paranodal gap (p = 0.122). 247 248 There were significant differences between experimental groups in the percentage of atypical nodal complexes (Figure 3c-e; F = 374.951, DF = 3, p = 0.0001). Atypical nodal complexes 249 were defined as either a heminode, characterised as a β -III tubulin+ area flanked by only one 250 251 Caspr+ area, or a single paranode, defined as a Caspr+ area not associated with a β -III tubulin+ area (Figure e) (Szymanski et al. 2013). Injury resulted in an increase in the 252 percentage of atypical nodal complexes in the injured, vehicle treated group compared to the 253 sham injured, vehicle treated group (p = 0.0001), which was significantly reduced by both 254 lomerizine + oxATP + YM872 (p = 0.0001) and lomerizine + BBG + YM872 (p = 0.0001). 255 256 However, both ion channel inhibitor combination groups still had significantly increased levels of atypical nodal complexes compared to the sham injured, vehicle treated group (p =257 258 0.0001). There was no significant difference between the two ion channel inhibitor 259 combinations in the number of atypical nodal complexes (p = 0.668).

260 Effects of ion channel inhibitor combinations on the optokinetic nystagmus reflex

261 The optokinetic nystagmus test of visual function, revealed significant differences in the

number of smooth pursuits and fast resets following injury and treatment (Figure 4a; smooth

263 pursuits F = 7.05, DF = 3, p = 0.001; fast resets F = 7.656, DF = 3, p = 0.001). Injury resulted

264 in a significant decrease in the numbers of both smooth pursuits and fast resets by the injured, vehicle treated group compared to the sham injured, vehicle treated group (p = 0.002, p =265 0.001 respectively). Animals treated with lomerizine + oxATP + YM872 made significantly 266 267 more smooth pursuits than the injured, vehicle treated group (p = 0.035), but fewer fast resets than the sham injured, vehicle treated group (p = 0.006). Animals treated with lomerizine + 268 BBG + YM872 made significantly more smooth pursuits (p = 0.002) and fast rests (p =269 0.039) than the injured, vehicle treated group; outcomes were not different from the sham 270 injured, vehicle treated group (p = 0.992, p = 0.284 respectively). There was no significant 271 272 difference between the two ion channel inhibitor combinations in the number of smooth pursuits (p=0.422) or fast resets (p=0.222). 273 274 To control for the length of the tracking motions, the time engaging in tracking behaviour 275 was also assessed, giving similar outcomes (Figure 4b; F = 11.458, DF = 3, p = 0.0001).

276 Injury resulted in a significantly lower proportion of time spent engaging in smooth pursuits

by the injured, vehicle treated group compared to the sham injured, vehicle treated group (p =

278 0.0001). Animals treated with lomerizine + oxATP + YM872 spent more time tracking than

the injured, vehicle group (p = 0.01), but still less time than the sham injured, vehicle treated

group (p = 0.045). Animals treated with lomerizine + BBG + YM872 also spent significantly

longer performing smooth pursuits than the injured, vehicle treated group (p = 0.0001), and

the time spent tracking was not significantly different to the sham injured, vehicle treated

group (p = 0.161). There was no significant difference between the two ion channel inhibitor

combinations in the time spent tracking (p = 0.601).

285

286 Discussion

287 The aim of this study was to determine if the combination of lomerizine + oxATP + YM872, shown to effectively limit functional loss associated with secondary degeneration of the optic 288 nerve (Savigni et al. 2013), would be as effective if oxATP was replaced with BBG, a more 289 290 clinically applicable P2X₇ receptor inhibitor that can cross the closed blood brain barrier. It 291 was found that the lomerizine + BBG + YM872 combination was as effective, or marginally more effective at the tested concentrations, than the lomerizine + oxATP + YM872 292 293 combination, at preserving node/paranode structure and visual function when delivered locally. However, neither therapeutic combination affected numbers of microglia and 294 295 macrophages, or the number of OPCs or oligodendrocytes. The data suggest an associative 296 relationship between preservation of myelin structure and maintenance of visual function 297 following injury.

298 The observed increase in nodal and paranodal lengths following partial optic nerve injury is in line with previous findings (Szymanski et al. 2013; O'Hare Doig et al. 2017) and is 299 suggestive of myelin retraction and a breakdown of the paranodal junction 300 (Arancibia- Carcamo and Attwell 2014). Increased P2X7 receptor activation on the myelin 301 sheath has also been associated with myelin degradation following injury, however the 302 underlying cellular mechanisms remain unclear (Matute 2008). The lomerizine + BBG + 303 YM872 combination restored nodal and paranodal structure to dimensions closer to the sham 304 305 control group than the lomerizine + oxATP + YM872 combination. The lomerizine + BBG + YM872 combination was the only treatment to not be different to the sham control group for 306 307 visual function. BBG is a more potent and selective antagonist of P2X7 receptors than oxATP 308 (Donnelly-Roberts and Jarvis 2007), which may explain this marginally greater efficacy of 309 the BBG containing combination in preserving myelin structure and visual function compared to the control groups in the current study. However, titration of doses of these 310 311 agents relative to their specific inhibitory concentrations for P2X7 receptors is required to

definitively compare efficacy of the two agents within the context of this treatment combination and injury model. We have previously established that locally delivered oxATP alone does not preserve node/paranode structure nor visual function in the partial optic nerve transection model (Savigni et al. 2013). The addition of lomerizine and YM872 to the combination is required for full beneficial effects (Savigni et al. 2013), emphasising that limiting excess Ca^{2+} flux through voltage gated calcium channels and Ca^{2+} permeable AMPA receptors is also important.

Following injury to the CNS, there is a high influx of Ca^{2+} into myelin *via* AMPA receptors 319 (Fowler et al. 2003), which results in increased Ca^{2+} binding to the catalytic core of calpain, 320 enhancing calpain activation (Croall and Demartino 1991; Khorchid and Ikura 2002). An 321 increase in calpain activation can induce myelin degradation, via cleavage of myelin basic 322 323 protein and myelin-associated glycoprotein (Banik et al. 1985; Shields et al. 1997; Fu et al. 2007). Myelin degradation has been associated with paranodal loop eversion and sheath 324 retraction and thus increased nodal and paranodal lengths (Ouyang et al. 2010). Therefore, 325 observed preservation of nodal and paranodal length by both ion channel combinations may 326 be due to inhibition of this Ca²⁺-dependent calpain mechanism through the antagonistic 327 activity of YM872 on AMPA receptors. 328

Furthermore, when the axolemma becomes exposed following myelin sheath retraction, and 329 paranodal splitting, there is an increase in Ca²⁺entry into axons *via* sub-myelin L-type 330 331 VGCCs, which are normally hidden underneath the myelin sheath (Zhang and David 2015). This contributes to neuronal Ca²⁺ overload, associated with oxidative stress, caspase-332 mediated apoptosis and decreased function (Annunziato et al. 2003). Lomerizine-mediated 333 inhibition of these sub-myelin VGCCs from beneath the sheath, together with exposure of 334 fewer L-type VGCCs by prevention of the myelin retraction, may be a further therapeutic 335 mechanism of the combinations of inhibitors. 336

337 Myelin structure is integral to the capacity of nerves to propagate action potentials, with the lengthening of the node associated with slower conduction velocities in a variety of 338 pathologies (Howell et al. 2006; Reimer et al. 2011; Sun et al. 2012). Abnormal myelination 339 340 at a single internode can be sufficient to block neural signal transduction for an entire axon (Baumann and Pham-Dinh 2001). Previous studies have also hypothesised that abnormalities 341 in the node of Ranvier proteins, associated with increased nodal length, may result in 342 343 decreased synchronicity of neuronal firing (Arancibia- Carcamo and Attwell 2014). 344 Preservation of myelin integrity by the ion channel inhibitor combinations may be facilitating 345 action potential propagation along axons, associated with preservation of function following injury. 346

347 However, in the current study myelin structure and visual function were only partly preserved by the ion channel inhibitor treatments, which suggests some aspects of myelin breakdown 348 following injury are mediated via alternative mechanisms of damage. One potential 349 350 mechanism is immune-cell mediated depletion of OPCs. The current study found that following injury, there is a significant increase in infiltrating microglia and macrophages, but 351 not resident microglial cells, indicative of an infiltrating immune response. However, 352 treatment did not show a significant effect at ameliorating this infiltrating immune response. 353 Following neurotrauma, inflammatory cells produce cytokines and chemokines, as well as 354 reactive oxygen species, resulting in oxidative damage of surrounding tissue (Anderson 355 2002). OPCs are especially vulnerable to oxidative stress and apoptosis following injury 356 (Thorburne and Juurlink 1996; Giacci et al. 2018), which may be why the combinations of 357 358 ion channel inhibitors were unable to ameliorate the loss of OPCs in this study. OPCs are required for oligodendrogenesis and remyelination following injury (Mirron et al. 2011). 359 OPCs also contribute to the formation of myelin nodal structures (Butt et al. 2004). It may be 360 361 that significantly preserving OPCs following injury, perhaps through preventing this

infiltrating immune response, would be associated with a complete preservation of myelin
structure and thus visual function. Furthermore, given previous studies have found an
increased therapeutic effect of the lomerizine + oxATP + YM872 combination after three
months of administration compared to three days (Savigni et al. 2013; O'Hare Doig et al.
2017), it may be that a longer duration of treatment would provide further improvements to
these outcomes following injury.

This study showed that the combination of lomerizine + BBG + YM872, which has the 368 potential to be delivered systemically following injury, shows promise for limiting secondary 369 370 degeneration following neurotrauma, however further work remains to be done. While BBG is a highly selective P2X7 receptor antagonist, it is 30 to 50 times more potent in rats than 371 humans (Jiang et al. 2000), which will necessitate careful titration of dosages before clinical 372 373 translation will be feasible. Lomerizine is currently used in clinical practice (Hara et al. 1999), YM872 is more soluble than other Ca^{2+} permeable AMPA receptor inhibitors 374 (Takahashi et al. 2002) and has been trialled in stroke with an acceptable safety profile 375 (Labiche and Grotta 2004) and BBG has no-observed-adverse-effect level of 8966mg/kg per 376 day in a mouse model of lifetime toxicity (Borzelleca et al. 1990). Nevertheless, while no 377 378 adverse effects have been observed in our rodent model, the combination of three ion channel 379 inhibitors will need to be careful assessed for toxicity in humans before a trial of efficacy 380 following neurotrauma can be contemplated. Furthermore, given the clinical need for 381 systemically administered drug delivery following neurotrauma, the efficacy of this blood brain barrier permeable combination of lomerizine + BBG + YM872 needs to be tested when 382 systemically delivered following injury. 383

384

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529 Figure Legends

Fig. 1 Effects of ion channel inhibitor combinations on densities of Iba1+ and ED1+ cells.

531 Densities of Iba1+(a), ED1+(b) and Iba1+/ED1+(c) cells in the ventral optic nerve from

sham injured, vehicle treated animals, injured, vehicle treated animals, and ion channel

inhibitor treated animals 3 days after partial optic nerve transection. N = 7 - 10 rats per group;

534 graphs display min to max values, with the central line representing the median data point.

535 Significant differences are indicated by * ($p \le 0.05$). (d) Representative images of Iba1+,

ED1+ and Iba1+/ED1+ cells, indicated with arrow heads; scale bar = $10\mu m$

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Fig. 2 Effects of ion channel inhibitor combinations on oligodendroglial cells. Densities of Olig2+ oligodendroglial cells (a) and PDGF α R+/Olig2+ OPCs (c) in the ventral optic nerve from sham injured, vehicle treated animals, injured, vehicle treated animals, and ion channel inhibitor treated animals, 3 days after partial optic nerve transection. N = 8 - 10 rats per group. Graphs display min to max values, with the central line representing the median data point. Significant differences are indicated by ** (p ≤ 0.01). Representative image of Olig2+ cells (b) and OPCs (d), indicated with arrow heads; scale bar = 15µm and 10µm respectively

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Fig. 3 Effect of ion channel inhibitor combinations on node/paranode complexes. Paranode length (a), paranodal gap length (b) and percentage of atypical node/paranode complexes (c) from thirty nodal complexes per animal in the ventral optic nerve from sham injured, vehicle treated animals, injured, vehicle treated animals, and ion channel inhibitor treated animals 3 days after partial optic nerve transection. Graphs display min to max values, with the central line representing the median data point; N = 6 - 9 rats per group. Significant differences indicated by * p \leq 0.05, ** p \leq 0.01, and *** p \leq 0.001. Differences compared to the sham injured, vehicle treated group are indicated by *, differences compared to the injured, vehicle treated group are indicated by #. (d) Representative image of two Caspr+ areas flanking a β -III tubulin+ area i.e. a typical nodal complex; scale bar = 2 μ m. (e) Representative image of one Caspr+ area flanking a β -III tubulin+ area denoting an atypical nodal complex / heminode; scale bar = 2 μ m

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559 Fig. 4 Effects of injury and ion channel inhibitor combinations on the number of responses in the optokinetic nystagmus test of visual function. Total number of smooth pursuits and fast 560 resets per minute engaged in the task (a) and proportion of time paying attention to task 561 engaged in smooth pursuits (b) by sham injured, vehicle treated animals; injured, vehicle 562 treated animals; or injured ion channel inhibitor treated animals. Graphs display min to max 563 564 values, with the central line representing the median data point; N = 5 - 10 rats per group. Significant differences are indicated by * $p \le 0.05$, ** $p \le 0.01$, and *** $p \le 0.001$. 565 Differences compared to the sham injured, vehicle treated group are indicated by *. 566 differences compared to the injured, vehicle treated group are indicated by # 567