

1 Transcriptomic Analysis of A Cannabis-Derived

2 Neuroprotective Therapy in a Zebrafish Model of ALS

3 AUTHORS AND AFFILIATIONS

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11 ABSTRACT

12 Amyotrophic lateral sclerosis (ALS) is a progressive neurodegenerative disease characterized by
13 motor neuron loss, and currently has limited therapeutic options. To advance the development of
14 effective interventions, we conducted a comprehensive transcriptomic analysis using a zebrafish
15 model of ALS induced by β -N-methylamino-L-alanine (BMAA) exposure. Zebrafish embryos
16 were treated with candidate neuroprotective agents, including CNR-401, Edaravone, and
17 Cannflavin A, and assessed for phenotypic and molecular responses. Leveraging Canurta's DEG

18 Pipeline Assistant, we performed high-throughput RNA-seq analysis, identifying robust
19 differential gene expression signatures and pathway enrichments associated with both disease
20 and therapeutic intervention. Our results demonstrate that BMAA exposure induces significant
21 motor deficits and widespread transcriptomic alterations, notably upregulating genes involved in
22 neuroinflammation and extracellular matrix (ECM) remodeling. Importantly, treatment with
23 CNR-401 not only produced the most extensive transcriptomic response but also significantly
24 rescued BMAA-induced motor deficits in zebrafish larvae, directly linking molecular changes to
25 phenotypic improvement. Functional enrichment analyses revealed that CNR-401 modulates
26 processes central to ALS pathology, including inflammatory response, ECM organization,
27 calcium signaling, steroid metabolism, and neuronal survival via the PI3K-Akt pathway.
28 Edaravone and Cannflavin A elicited more limited and distinct transcriptomic changes, with
29 Edaravone primarily affecting metabolic and signaling pathways, and Cannflavin A modulating
30 metabolic and steroidogenic functions.

31 INTRODUCTION

32 ALS is a rapidly progressive neurodegenerative disease characterized by the loss of motor
33 neurons, leading to muscle weakness, paralysis, and ultimately death. Despite decades of
34 research, effective treatments remain scarce, and the disease burden continues to grow globally.
35 Existing therapies, such as riluzole and edaravone, offer only modest benefits [1], underscoring
36 the urgent need for new therapeutic strategies that target the complex molecular mechanisms
37 underlying ALS pathogenesis.

38 Zebrafish (*Danio rerio*) provide a powerful vertebrate model for studying neurodegenerative
39 diseases due to their genetic tractability, conserved neurobiology, and suitability for

40 high-throughput screening. The neurotoxin BMAA is reliably known to induce ALS-like
41 phenotypes in zebrafish, enabling the evaluation of candidate neuroprotective compounds in vivo
42 [2].

43 Canurta Therapeutics is dedicated to advancing the development of novel ALS therapies through
44 rigorous preclinical research and innovative computational tools. This study presents the first use
45 of the newly developed DEG Pipeline Assistant that facilitates reproducible, high-throughput
46 analysis of RNA-seq data, enabling rapid identification of disease-relevant genes and pathways
47 that help inform therapeutic strategies [3].

48 BMAA-Treated Zebrafish Model of ALS

49 To model ALS-like neurodegeneration, we utilized β -methylamino-L-alanine (BMAA), a
50 cyanobacterial neurotoxin known to induce motor neuron toxicity and behavioral deficits in
51 zebrafish [2]. Neurotoxins, and specifically BMAA, are increasingly implicated in ALS
52 pathogenesis [4]. Dietary exposure to BMAA through bioaccumulation in aquatic food webs has
53 been directly linked to ALS clusters, such as the 50-100-fold increase in
54 ALS-parkinsonism-dementia complex observed among the Chamorro people of Guam who show
55 an average of 6.6 $\mu\text{g/g}$ BMAA in their brain tissues [5]. BMAA is thought to cause this
56 neurodegeneration through pathways such as microglial activation, chronic inflammation, and
57 calcium overload [6].

58 Exposure to BMAA in zebrafish reliably induces ALS-like phenotypes including motor deficits,
59 spinal abnormalities, altered neuromuscular junction morphology, and increased oxidative stress
60 [2], enabling the evaluation of candidate neuroprotective compounds in vivo. Zebrafish share

61 approximately 70% of their genes with humans, and an estimated 84% of human
62 disease-associated genes have at least one zebrafish ortholog [7], making them a genetically
63 tractable system for modeling human pathologies. Their nervous system exhibits conserved
64 neuroanatomy and neurochemistry, including orthologous cannabinoid receptors (CB1 and CB2)
65 [8], which are critical for evaluating cannabinoid and cannflavin-based therapeutics.
66 Additionally, zebrafish embryos develop rapidly, are optically transparent, and are amenable to
67 high-throughput screening, enabling efficient assessment of neurobehavioral phenotypes and
68 drug efficacy. This approach leverages the genetic, physiological, and pharmacological
69 similarities between zebrafish and humans, providing a robust platform for the preclinical
70 assessment of ALS therapeutics targeting cannabinoid pathways.

71 CNR-401: A Novel Cannabis-Derived Formulation

72 CNR-401 represents a strategically assembled combination of bioactive compounds selected for
73 their established neuroprotective and anti-inflammatory properties. This approach leverages the
74 therapeutic potential of phytochemicals derived from *Cannabis sativa* to address pathological
75 changes associated with neurodegenerative conditions. Among other primary cannabinoids,
76 cannabidiol (CBD) serves as the primary active component in this formulation, demonstrating
77 extensive neuroprotective properties through anti-inflammatory and antioxidant effects [9]. CBD
78 has generated significant interest in its therapeutic potential against secondary injury cascades in
79 traumatic brain injuries, which share key pathological characteristics of ALS such as glial cell
80 dysregulation, excitotoxicity, and neuroinflammation [10]. The inclusion of cannabinoid acid
81 precursors further broadens the therapeutic spectrum of CNR-401, as these molecules are known
82 to offer distinct benefits compared to their decarboxylated counterparts [11]. The formulation is
83 also enhanced by select terpene ingredients, which contribute additional anti-inflammatory and

neuroprotective actions [12]. Cannflavin A, a unique prenylated flavonoid, is incorporated due to its exceptional anti-inflammatory properties achieved through dual inhibition of microsomal prostaglandin E2 synthase-1 (mPGES-1) and 5-lipoxygenase (5-LOX) [13], having been found to be ~30 times more effective than aspirin in reducing inflammation [14]. The strategic design of CNR-401 leverages the entourage effect [15,16] in which cannabinoids, terpenes, and flavonoids interact synergistically to amplify overall therapeutic effectiveness. To comprehensively assess its neuroprotective potential in ALS, this study directly compared CNR-401's transcriptomic effects with those of a clinically approved positive control and pure Cannflavin A within the zebrafish model.

MATERIALS AND METHODS

Solvent Toxicity Assessment

Solubilizers were necessary to make cannabinoids and terpenes in the test formulations miscible in water, as these compounds are inherently not water-soluble, creating challenges for aquatic organism testing. Two cyclodextrin solubilizers were evaluated: hydroxypropyl- β -cyclodextrin (HPC) and hydroxyethyl- β -cyclodextrin (HEC). Cyclodextrins are cyclic oligosaccharides that act as solubilizing agents and are utilized in many commercially available drugs.

For each cyclodextrin, 100 mM stock solutions in water were prepared. Toxicity assays for both cyclodextrins were completed at 6 days post-treatment and performed at concentrations of 0, 0.5, 1, 5, 10, and 20 mM. Following positive results with HEC, extended toxicity testing was conducted with concentrations of 0, 15, 20, 30, 40, and 50 mM using a freshly prepared stock solution. Based on these results, all subsequent experiments were performed using 10 mM HEC

105 as the final concentration to maintain safety margins while accounting for potential additive
106 toxicity from test compounds.

107 Product Toxicity Assessment

108 Toxicity assessments were performed on CNR-401 and Cannflavin A at final concentrations of 0,
109 0.5, 1, 2, 5, and 10 μ M. CNR-401 concentrations were expressed in terms of the highest
110 concentration principal active component (CBD) of the complex botanical mixture. Cannflavin A
111 was used as a reference standard. All toxicity assays were completed at 6 days post-treatment
112 with HEC maintained at 10 mM final concentration in all conditions. Diluted solutions were
113 prepared in HEC 5-10 minutes before addition to multi-well plates.

114 Product Efficacy Assessment and Phenotypic Analysis

115 At 8 hours post-fertilization, zebrafish embryos were distributed into 48-well plates (1 per well)
116 and exposed to BMAA at optimized concentrations to induce ALS-like neurodegeneration. Test
117 compounds (CNR-401 and Cannflavin A) were administered simultaneously at their respective
118 NOTEC concentrations. Each experimental plate contained non-treated controls, BMAA-treated
119 controls, and BMAA + compound-treated wells (8 wells per condition). Edaravone, the currently
120 available ALS medication, was included as a positive control. Plates were incubated at 28°C
121 throughout the experimental period.

122 Behavioral assessments were conducted using the DanioVision automated behavioral tracking
123 system [17] equipped with a GigE camera. At 6 days post-fertilization, zebrafish larvae were
124 placed in the DanioVision system and exposed to white light (15–16 μ M/s/m) for 1 minute,
125 followed by tracking for 20 minutes under light conditions. Locomotion patterns were recorded

126 and analyzed using EthoVision XT17 software [17] to generate comprehensive behavioral
127 profiles. Following behavioral recording, larvae were visually examined to assess phenotypic
128 responses and remove any dead, necrotic, or morphologically affected specimens from the
129 analysis. Visual assessments included evaluation of cardiac edema, abnormal heart rate, body
130 deformation, and overall morphological integrity. Only larvae displaying normal morphology
131 and survival were included in the final behavioral analysis to ensure data quality and
132 experimental validity. Distance moved (mm) served as the primary behavioral endpoint, with
133 BMAA treatment expected to significantly reduce locomotion compared to untreated controls.
134 Effective neuroprotective compounds were anticipated to demonstrate rescue effects by restoring
135 locomotor activity toward control levels.

136 Experimental Conditions and Replication

137 To investigate transcriptomic responses to neuroprotective therapies in a zebrafish model of
138 ALS, we designed a multi-condition experiment utilizing six biological replicates per group.
139 Zebrafish embryos were randomly assigned to one of five experimental conditions:

- 140 ● Control: Untreated zebrafish embryos, serving as the baseline reference
- 141 ● BMAA: Embryos exposed to β -N-methylamino-L-alanine to induce neurodegenerative
142 phenotypes characteristic of ALS
- 143 ● CNR-401: BMAA-exposed embryos subsequently treated with Canurta's drug candidate
144 CNR-401
- 145 ● Edaravone: BMAA-exposed embryos subsequently treated with edaravone, a clinically
146 approved ALS therapy
- 147 ● Cannflavin A: BMAA-exposed embryos subsequently treated with cannflavin A

148 Sample Preparation and RNA Sequencing

149 Non-control zebrafish embryos were exposed to BMAA to induce neurodegeneration, with
150 additional groups receiving candidate neuroprotective agents, including edaravone and
151 CNR-401. RNA was extracted using the Qiagen RNeasy kit and libraries were prepared with the
152 Illumina TruSeq Stranded RNA kit. Sequencing was performed on an Illumina NovaSeq 6000
153 platform, yielding an average of 36.3 million read pairs per sample.

154 Data Processing and Visualization

155 Raw reads were trimmed with cutadapt [18] and mapped to the Danio rerio reference genome
156 using STAR [19,20]. Fastqc [21] was used to analyze the quality of the reads before and after
157 trimming. Gene-level counts were subsequently imported into DESeq2 [22] for normalization
158 and pairwise differential expression analysis. Lastly, the Canurta DEG Pipeline Assistant [3] was
159 employed for downstream analysis:

- 160 • Automated metadata construction and quality control (boxplots, correlation heatmap,
161 PCA)
- 162 • Differential expression filtering with \log_2 fold change > 1 and adjusted p-value < 0.05
- 163 • Visualization (MA plots, volcano plots, heatmaps)
- 164 • Ortholog mapping to human genes
- 165 • GO and KEGG functional enrichment analysis with adjusted p-value < 0.05

166 (Supplementary Material: Screenshot of DEG Pipeline Assistant Parameters Used).

167 RESULTS

168 Solvent Toxicity Assessment

169 HPC demonstrated significant toxicity at 5 mM concentration, resulting in 75% larval mortality
170 with observable toxicity in surviving larvae including cardiac edema, abnormal heart rate, and
171 body deformation. Complete mortality was observed at 10 mM and 20 mM concentrations. In
172 contrast, HEC showed no toxicity up to 20 mM, with all larvae surviving and displaying no
173 morphological changes or significant decrease in movement.

174 Product Toxicity Assessment

175 Both CNR-401 and Cannflavin A demonstrated no toxicity up to 2 μ M concentration
176 (Supplementary Material: Product Toxicity Assays). CNR-401 showed 62.5% larval survival at 5
177 μ M and complete mortality at 10 μ M, yielding an LC50 of 4.827 μ M. Cannflavin A exhibited
178 37.5% survival at 5 μ M and complete mortality at 10 μ M, with an LC50 of 4.694 μ M.

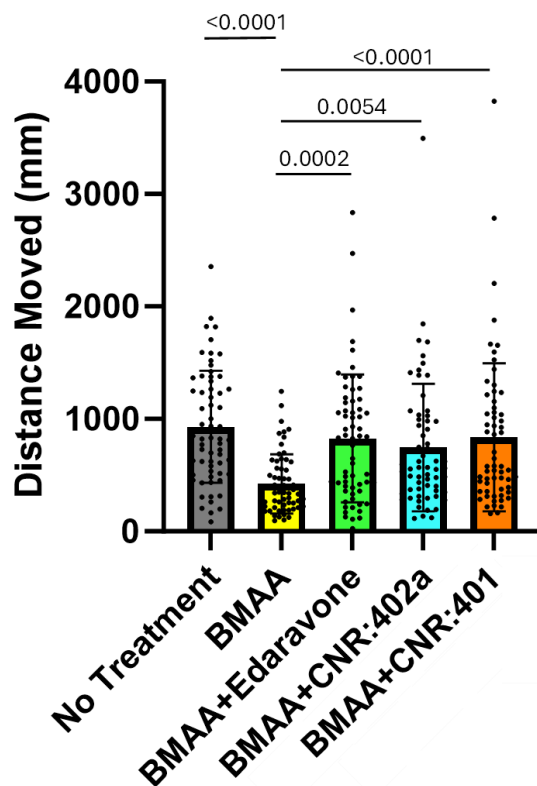
179 Teratotoxicity assessments closely followed acute toxicity trends for both compounds.

180 Given that both products were safe up to a final concentration of 2 μ M, the assays were repeated
181 in three biological experiments in order to explore the toxicity between 2 and 5 μ M to find the
182 maximum safe concentrations.

183 Product Efficacy Assessment

184 BMAA treatment significantly reduced locomotion in zebrafish larvae ($p < 0.0001$), confirming
185 the validity of the neurodegeneration model. Edaravone, the current ALS medication,
186 demonstrated significant rescue effects ($p = 0.0002$) on BMAA-treated larvae mobility. Both
187 CNR-401 and Cannflavin A showed significant rescue effects with p-values of $p < 0.0001$ and p

188 = 0.0054, respectively (Figure 1). All treatments were administered at 2 μ M concentration with
 189 nine biological replicates per condition.



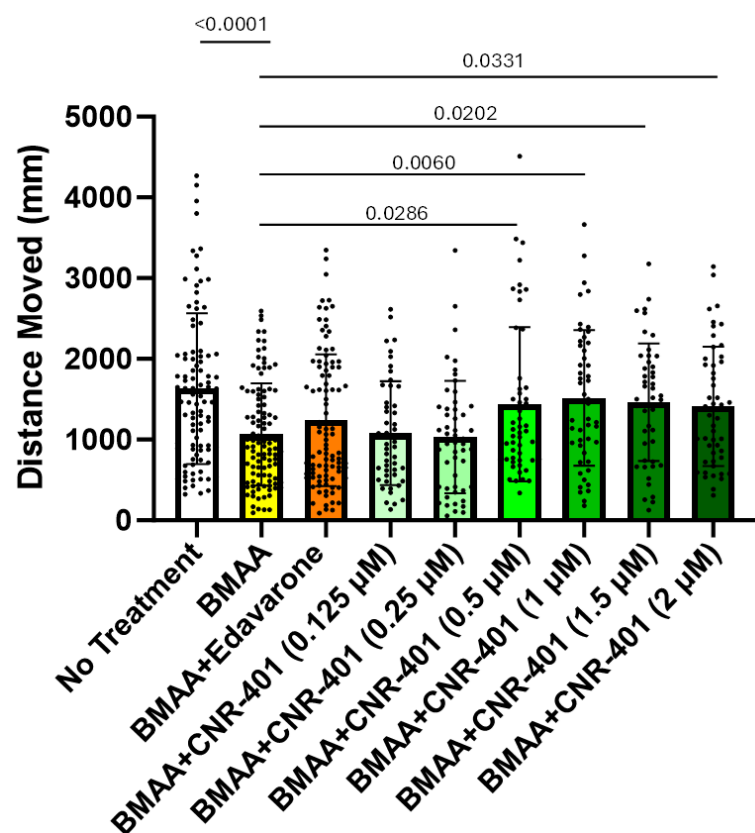
190

191 **Figure 1.** Effect of CNR-401 and Cannflavin A (CNR-402a) in a BMAA-induced model of
 192 neurotoxicity in zebrafish larvae. All treatments are dosed at 2 μ M. Experiments were done with
 193 nine biological replicates.

194 Product Dose-Response Analysis

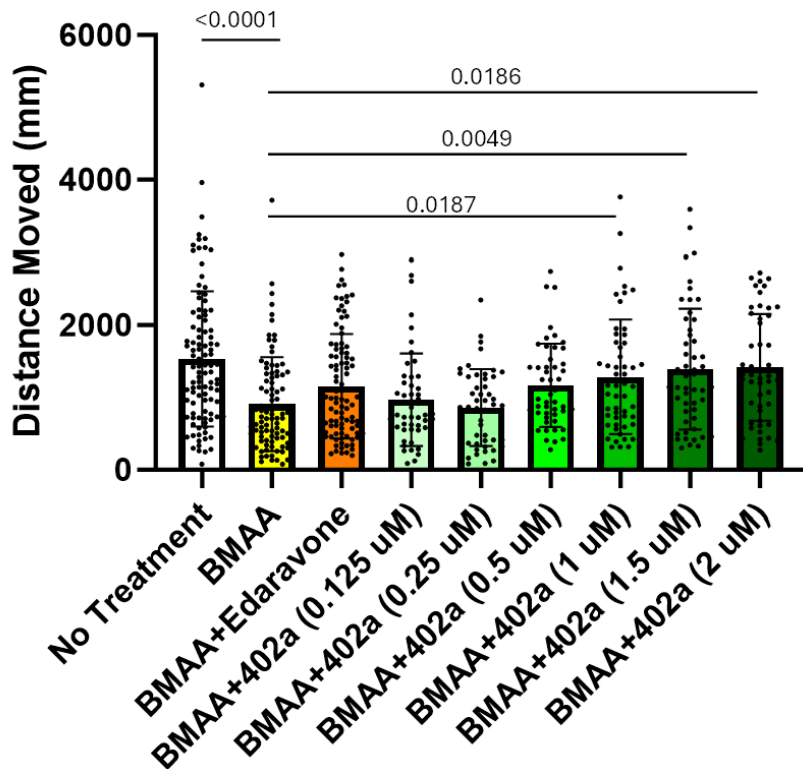
195 CNR-401 demonstrated dose-dependent efficacy with the lowest effective dose of 0.5 μ M in the
 196 BMAA-induced neurotoxicity model (Figure 2). Cannflavin A required a higher concentration,
 197 showing efficacy beginning at 1 μ M (Figure 3). This difference indicates superior potency for

198 CNR-401 at lower concentrations. Seven biological replicates were completed for each condition
 199 in the dose-response studies.



200

201 **Figure 2.** Dose response of CNR-401 in a BMAA-induced model of neurotoxicity in zebrafish
 202 larvae. Experiments were done with seven biological replicates.



203

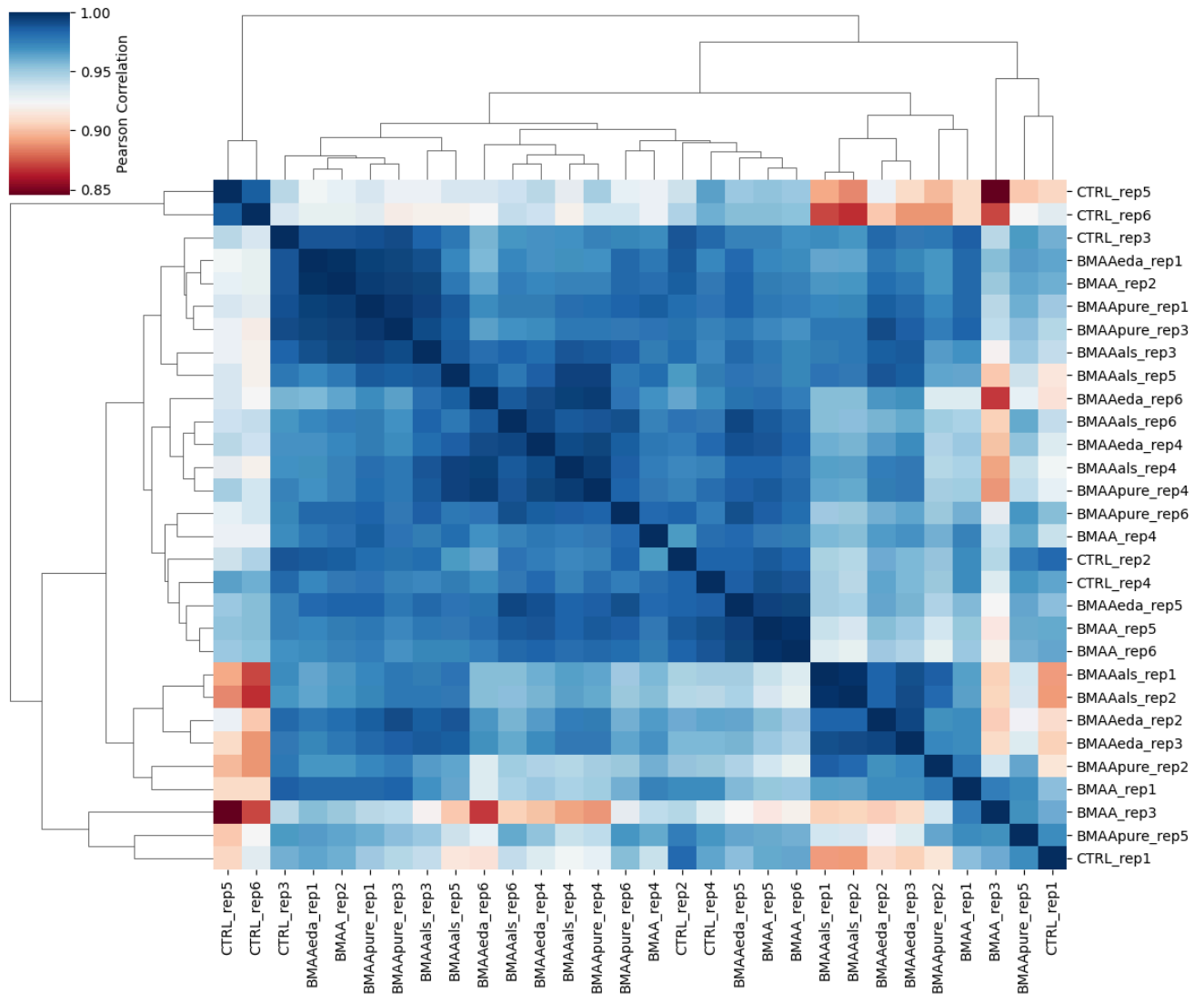
204 **Figure 3.** Dose response of Cannflavin A (402a) in a BMAA-induced model of neurotoxicity in
 205 zebrafish larvae. Experiments were done with seven biological replicates.

206 Data Quality

207 Quality control assessment through boxplot visualization (Supplementary Material: Plots)
 208 confirmed excellent data consistency across the biological replicates within each condition
 209 group. The normalized gene expression counts displayed remarkably uniform distribution
 210 patterns, with median expression levels precisely aligned across all six replicates for each
 211 condition, indicating successful normalization and comparable library sizes. The characteristic
 212 compression of interquartile ranges near zero reflects the expected transcriptomic landscape
 213 where the majority of genes exhibit low to moderate expression levels. Consistent outlier
 214 patterns extending to approximately 500,000 normalized counts were observed across all

215 samples, representing highly expressed genes such as housekeeping proteins that were
216 reproducibly detected. This uniform distribution profile, combined with the absence of
217 sample-specific technical artifacts or batch effects, demonstrates robust experimental
218 reproducibility and validates the dataset's suitability for downstream differential expression
219 analysis. The boxplot patterns align with established quality standards for normalized RNA-seq
220 data, confirming that technical variation has been effectively minimized while preserving
221 genuine biological signals.

222 The sample correlation analysis (Figure 4) demonstrated high reproducibility within
223 experimental groups, with control samples showing strong internal correlations ($r > 0.90$).
224 Treatment groups exhibited distinct expression profiles compared to controls, as evidenced by
225 the hierarchical clustering pattern that segregated samples by treatment condition. The overall
226 correlation structure supports the validity of the experimental design and suggests that the
227 observed gene expression changes are treatment-specific rather than due to technical variability.



228

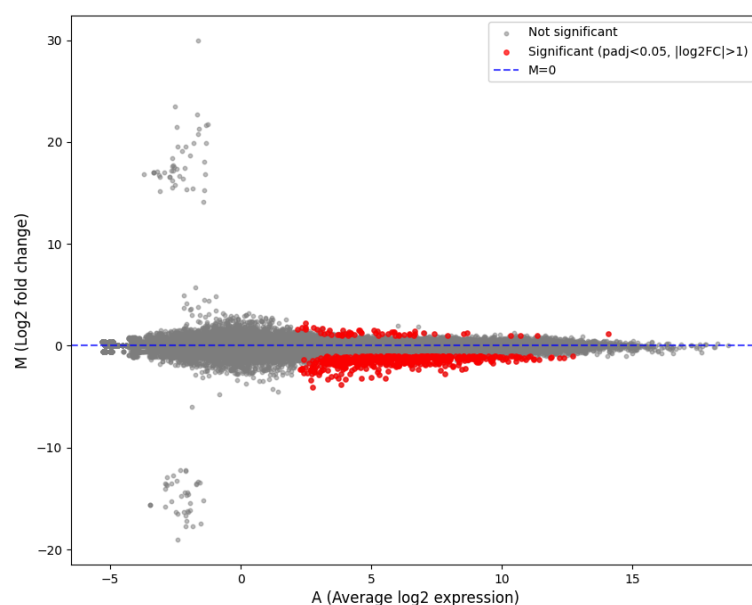
229 **Figure 4.** Heatmap showing pairwise Pearson correlations of gene expression between all
 230 samples, clustered hierarchically. A darker blue square indicates a stronger positive correlation,
 231 as seen in the squares along the diagonal showing a perfect self-correlation ($r=1.0$).

232 The principal component analysis (Supplementary Material: Plots) revealed substantial
 233 variability in gene expression profiles across experimental conditions. Notably, control samples
 234 showed considerable dispersion across the PCA space, with replicates distributed from the
 235 bottom region ($PC1 \geq 150$, $PC2 \approx -150$) to the upper right quadrant ($PC1 \approx 150$, $PC2 \approx 50$),

236 indicating significant baseline heterogeneity in untreated samples. BMAA treatment and
237 therapeutic interventions (CNR401, Edaravone, and Cannflavin A) showed similarly dispersed
238 patterns across both principal components, which together explain 38.04% of the total variance.
239 This observed variability in RNA-seq data aligns with established norms [23].

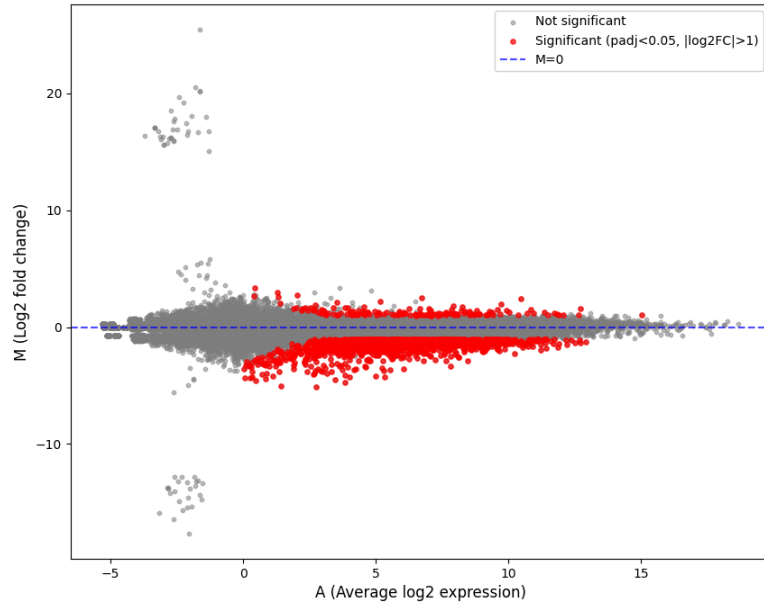
240 Differentially Expressed Genes

241 Differential gene expression analysis between BMAA-affected and treated conditions revealed
242 distinct therapeutic responses across the three neuroprotective compounds (Figures 5-6).
243 CNR-401 treatment resulted in 1,576 significantly differentially expressed genes, while
244 Edaravone treatment showed 359 differentially expressed genes, and Cannflavin A treatment
245 demonstrated 130 differentially expressed genes (Table 1). These findings indicate varying
246 degrees of transcriptomic modulation, with CNR-401 exhibiting the most extensive gene
247 expression changes relative to BMAA-affected conditions.



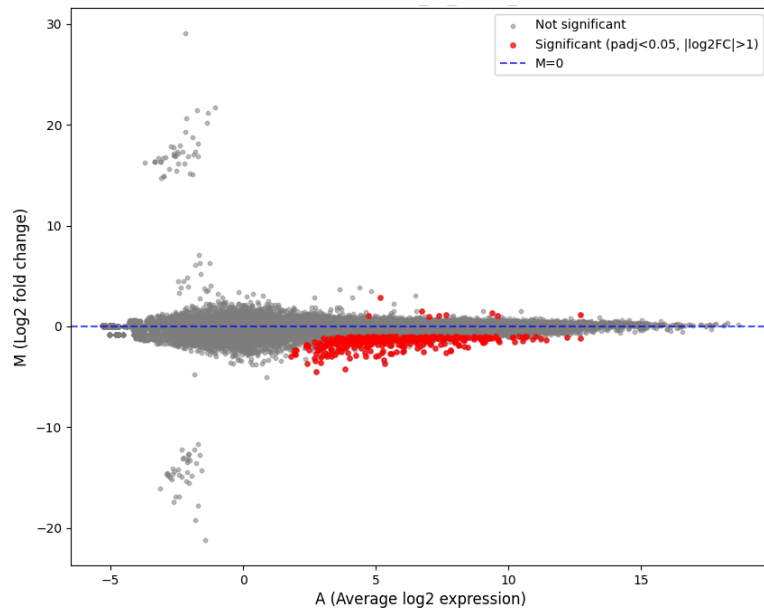
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249 **(a)** BMAA-Affected vs Control



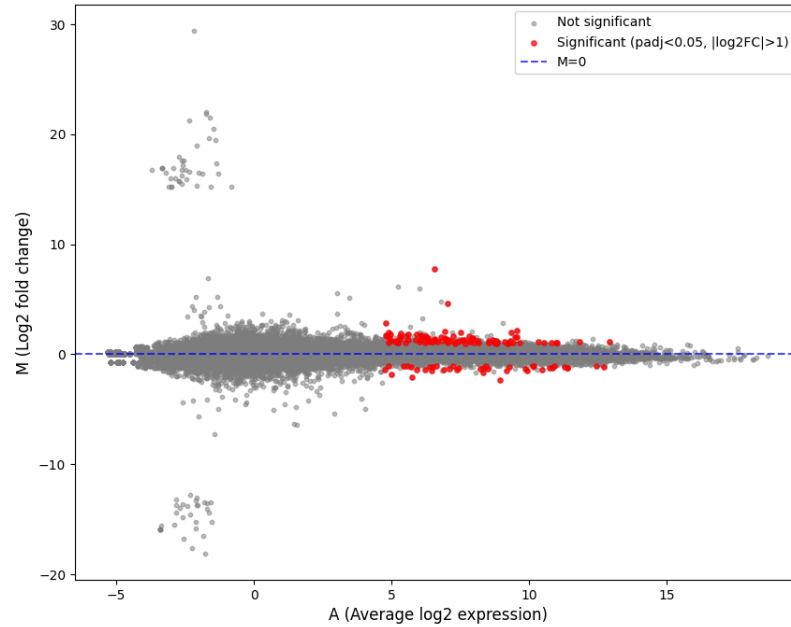
250

251 **(b)** BMAA-Affected vs CNR-401-Treated



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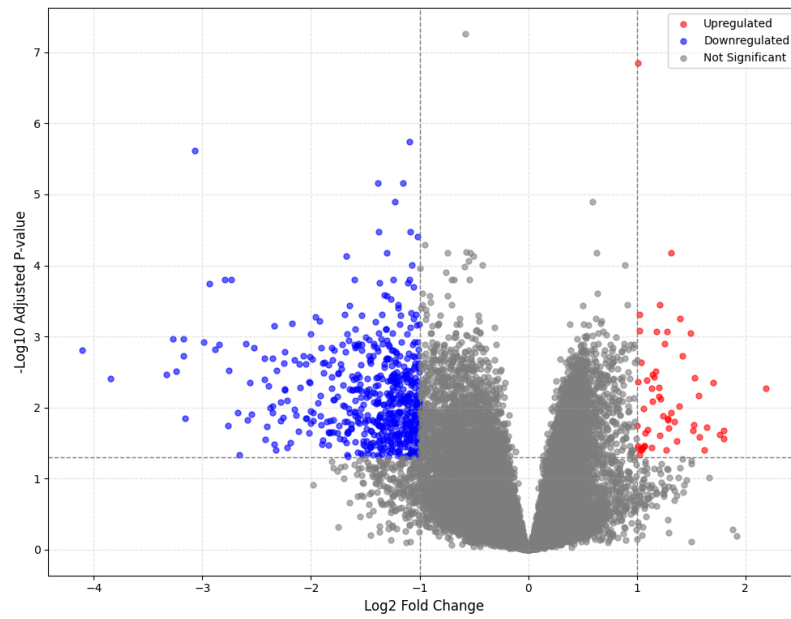
253 **(c)** BMAA-Affected vs Edaravone-Treated



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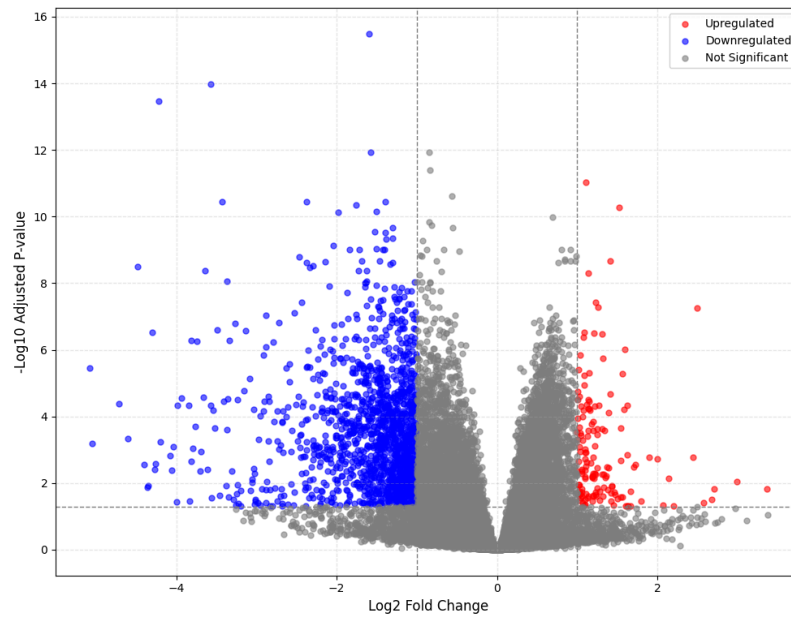
255 **(d)** BMAA-Affected vs Cannflavin A-Treated

256 **Figure 5.** MA plots showing differential gene expression analysis between BMAA-affected and
 257 treated conditions. Each point represents a gene plotted by average log expression (A, x-axis)
 258 versus log2 fold change (M, y-axis). Red points indicate significant differentially expressed
 259 genes.



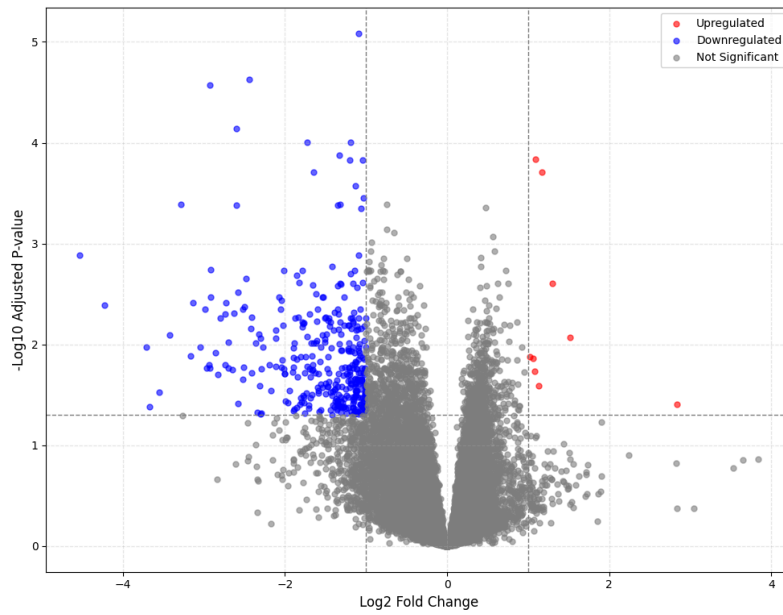
260

261 **(a)** BMAA-Affected vs Control



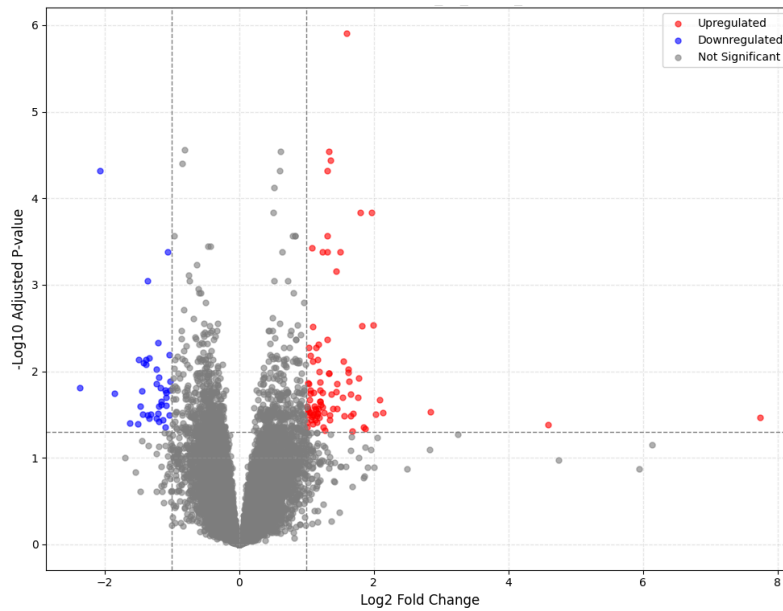
262

263 **(b)** BMAA-Affected vs CNR-401-Treated



264

265 (c) BMAA-Affected vs Edaravone-Treated



266

267 (d) BMAA-Affected vs Cannflavin A-Treated

268 **Figure 6.** Volcano plots displaying statistical significance versus biological significance of
 269 differential gene expression. Each point represents a gene plotted by log2 fold change (x-axis)

versus -log10 adjusted p-value (y-axis). Blue points indicate significantly downregulated genes, red points indicate significantly upregulated genes.

Table 1. Summary of pairwise comparisons of interest.

Comparison	Total Genes	Significant Genes	Upregulated	Downregulated
BMAA vs Control	28788	624	57	567
BMAA vs BMAA + CNR401	28788	1576	149	1427
BMAA vs BMAA + Edaravone	28788	359	9	350
BMAA vs BMAA + Cannflavin A	28788	130	91	39

Ortholog Mapping

Mapping the significantly expressed DEGs from the BMAA-affected vs CNR-401-treated comparison revealed that the top 19 genes belonged to the *UGT* Gene Family and were all downregulated (Table 2). It is important to note that these 19 human UGT genes mapped from the same underlying zebrafish gene transcripts, explaining the identical or nearly identical expression values. This ortholog mapping thus indicates a coordinated downregulation of the *UGT* family rather than independent measurements. In the BMAA vs control comparison, this family of orthologs showed only approximately a -0.5 log2foldchange (Supplementary Material: Data), demonstrating the significant impact of CNR-401 on these pathways.

Table 2. The top 30 human orthologs (sorted by L2FC) differentially expressed when BMAA-affected samples were treated with CNR401.

Ortholog Name	Description	Absolute L2FC	Expression	Adjusted P-Value
UGT2B10	UDP glucuronosyltransferase family 2 member B10	4.077694	Downregulated	0.001473
UGT2A3	UDP glucuronosyltransferase family 2 member A3	4.077694	Downregulated	0.001473
UGT2B28	UDP glucuronosyltransferase family 2 member B28	4.077694	Downregulated	0.001473
UGT3A1	UDP glycosyltransferase family 3 member A1	4.077694	Downregulated	0.001473
UGT2B4	UDP glucuronosyltransferase family 2 member B4	4.077694	Downregulated	0.001473
UGT1A6	UDP glucuronosyltransferase family 1 member A6	4.077694	Downregulated	0.001473

UGT2B7	UDP glucuronosyltransferase family 2 member B7	4.077694	Downregulated	0.001473
UGT2A1	UDP glucuronosyltransferase family 2 member A1 complex locus	4.077694	Downregulated	0.001473
UGT2B15	UDP glucuronosyltransferase family 2 member B15	4.077694	Downregulated	0.001473
UGT2B17	UDP glucuronosyltransferase family 2 member B17	4.077694	Downregulated	0.001473
UGT2B11	UDP glucuronosyltransferase family 2 member B11	4.077694	Downregulated	0.001473
UGT1A9	UDP glucuronosyltransferase family 1 member A9	4.077694	Downregulated	0.001473
UGT1A8	UDP glucuronosyltransferase family 1 member A8	4.077694	Downregulated	0.001473

UGT1A10	UDP glucuronosyltransferase family 1 member A10	4.077694	Downregulated	0.001473
UGT1A7	UDP glucuronosyltransferase family 1 member A7	4.077694	Downregulated	0.001473
UGT1A4	UDP glucuronosyltransferase family 1 member A4	4.077694	Downregulated	0.001473
UGT2A2	UDP glucuronosyltransferase family 2 member A2	4.077694	Downregulated	0.001473
UGT1A3	UDP glucuronosyltransferase family 1 member A3	4.077694	Downregulated	0.001473
UGT1A5	UDP glucuronosyltransferase family 1 member A5	4.077694	Downregulated	0.001473
URGCP	upregulator of cell proliferation	3.702947	Downregulated	0.004319
DAO	D-amino acid oxidase	3.68739	Downregulated	0.001132

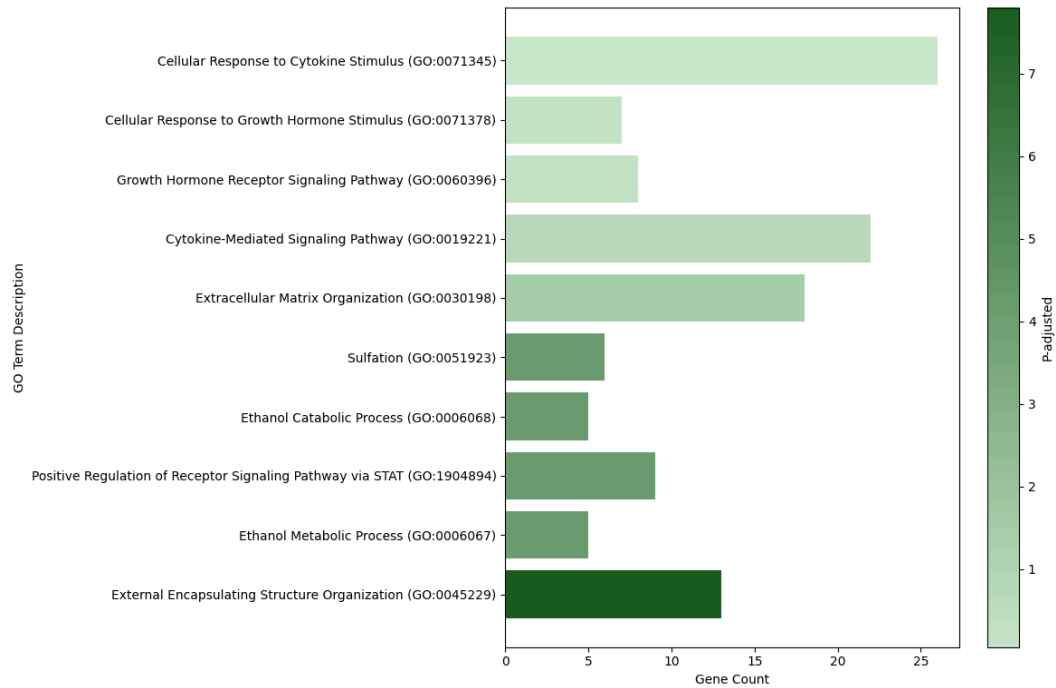
SDR42E2	short chain dehydrogenase/reductase family 42E; member 2	3.663107	Downregulated	2.679562886399 305e-05
GBP6	guanylate binding protein family member 6	3.558249	Downregulated	0.028347
MXRA5	matrix remodeling associated 5	3.424166	Downregulated	3.619114816823 018e-11
NOTCH4	notch receptor 4	3.15797	Downregulated	1.658254919506 955e-05
B3GALT9	beta-1;3-galactosyltransferase 9	3.018835	Downregulated	0.005205
HLA-DQA1	major histocompatibility complex; class II; DQ alpha 1	3.016801	Downregulated	0.034057
HLA-DQA2	major histocompatibility complex; class II; DQ alpha 2	3.016801	Downregulated	0.034057
UNG	uracil DNA glycosylase	2.996153	Upregulated	0.009045

COL6A3	collagen type VI alpha 3 chain	2.950076	Downregulated	9.900582448395 986e-05
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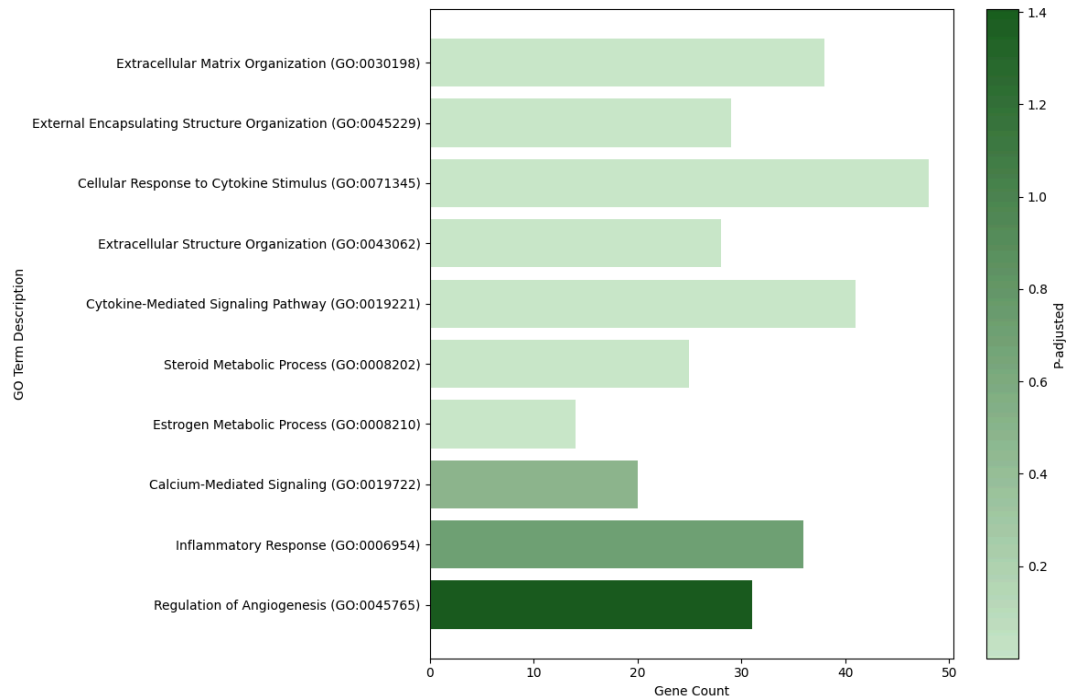
285 Functional Enrichments

286 Gene ontology (GO) biological process enrichment analysis revealed that, in the BMAA versus
287 BMAA+CNR401 comparison (Figure 7b), the most significantly enriched terms included
288 extracellular matrix organization, external encapsulating structure organization, cellular response
289 to cytokine stimulus, extracellular structure organization, cytokine-mediated signaling pathway,
290 steroid metabolic process, estrogen metabolic process, calcium-mediated signaling,
291 inflammatory response, and regulation of angiogenesis. For the BMAA versus control
292 comparison (Figure 7a), enriched GO biological processes included cellular response to cytokine
293 stimulus, cellular response to growth hormone stimulus, growth hormone receptor signaling
294 pathway, cytokine-mediated signaling pathway, extracellular matrix organization, sulfation,
295 ethanol catabolic process, positive regulation of receptor signaling pathway via STAT, ethanol
296 metabolic process, and external encapsulating structure organization.



297

298 (a) BMAA-Affected vs Control

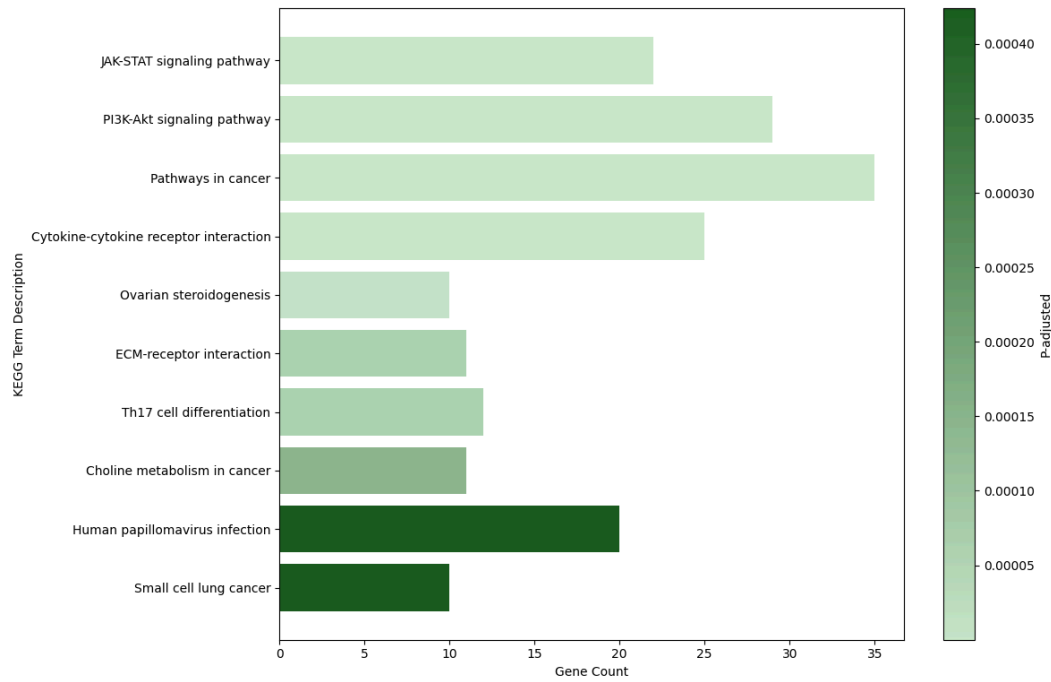


299

300 (b) BMAA-Affected vs CNR-401 Treated

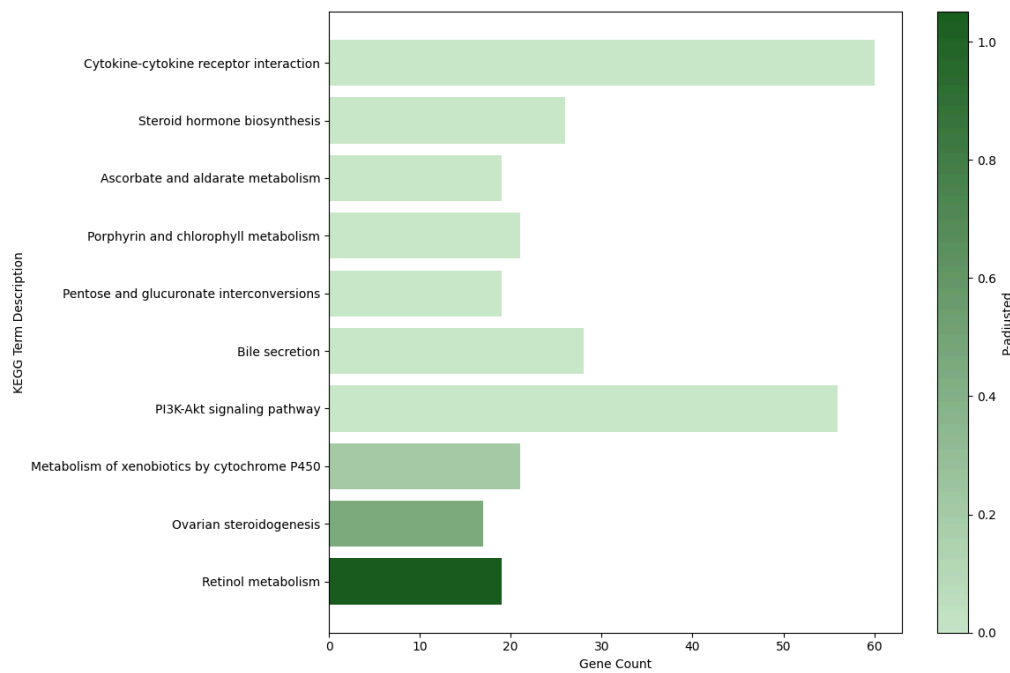
Figure 7. GO biological process (BP) enrichment analysis of differentially expressed genes between pairwise comparisons of interest. Bar length indicates the number of genes associated with each enriched GO term, while color intensity reflects the adjusted p-value for enrichment significance.

Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis showed that, in the BMAA versus BMAA+CNR401 comparison (Figure 8b), the top enriched pathways were cytokine-cytokine receptor interaction, steroid hormone biosynthesis, ascorbate and aldarate metabolism, porphyrin and chlorophyll metabolism, pentose and glucuronate interconversions, bile secretion, PI3K-Akt signaling pathway, metabolism of xenobiotics by cytochrome P450, ovarian steroidogenesis, and retinol metabolism. In the BMAA versus control comparison (Figure 8a), enriched KEGG pathways included JAK-STAT signaling pathway, PI3K-Akt signaling pathway, pathways in cancer, cytokine-cytokine receptor interaction, ovarian steroidogenesis, ECM-receptor interaction, Th17 cell differentiation, choline metabolism in cancer, human papillomavirus infection, and small cell lung cancer.



315

316 (a) BMAA-Affected vs Control



317

318 (b) BMAA-Affected vs CNR-401-Treated

Figure 8. KEGG pathway enrichment analysis of differentially expressed genes in each experimental comparison. Bar length represents the number of genes associated with each enriched pathway, and color intensity reflects the adjusted p-value for statistical significance.

DISCUSSION

CNR-401 Induces Broad Transcriptomic Changes in BMAA-Exposed Zebrafish

Our analysis revealed that CNR-401 treatment produces a strikingly extensive alteration in gene expression profiles compared to BMAA-alone exposure or subsequent treatment with other therapies. In the BMAA vs. BMAA + CNR-401 comparison, 1,576 genes were significantly differentially expressed, far exceeding the 359 DEGs observed with Edaravone and 130 with Cannflavin A. This indicates that CNR-401 elicits a broad transcriptomic response in the ALS model. Notably, the vast majority of CNR-401–modulated genes (1427 genes) were downregulated relative to BMAA-alone, with only 149 upregulated. This pattern suggests that CNR-401 largely counteracts or normalizes many of the gene expression changes induced by BMAA. In other words, genes pathologically elevated in the BMAA condition tend to be suppressed by CNR-401, pointing to a robust mitigation of BMAA-driven molecular perturbations. Such a widespread reversal of BMAA effects implies that CNR-401 engages multiple biological pathways, consistent with a multi-target mechanism of neuroprotection. In contrast, Edaravone and Cannflavin A triggered comparatively modest transcriptomic shifts, highlighting that CNR-401’s impact is both broader and more potent in this model. The fact that CNR-401 rescued diseased zebrafish in the functional mobility analysis at least confirms that this breadth of gene regulation is not harmful and instead may translate into a more comprehensive

therapeutic effect, as ALS pathology involves myriad dysregulated processes. However, this large amount of genetic changes does warrant careful interpretation.

Top Differentially Expressed Genes Reveal CNR-401's Multi-Target Neuroprotective Mechanisms

Several of the most dysregulated genes in the CNR-401 treatment group have clear links to ALS pathophysiology. *UDP-glucuronosyltransferase (UGT)* family members dominated the top-ranked DEGs. UGTs are Phase II detoxification enzymes responsible for glucuronidation of endogenous toxins and xenobiotics [25]. One explanation of their coordinated suppression suggests that CNR-401 reduced the need for BMAA detoxification via glucuronidation. In this interpretation, CNR-401 aids in neutralizing or clearing BMAA (or its toxic metabolites) by other means, thereby reducing reliance on endogenous UGT-mediated glucuronidation. In essence, if BMAA exposure alone induces *UGT* expression as a compensatory detox response, the addition of CNR-401 appears to alleviate that stress, resulting in lower UGT levels. A second interpretation supported by the functional enrichments is that the suppression of *UGT* genes reduces their known conjugation and elimination of steroids, thereby increasing levels of endogenous neuroprotective steroids, discussed later.

Several other ALS-relevant genes were significantly downregulated by CNR-401, aligning with beneficial modulation of neuroinflammatory and neurodegenerative pathways. *URGCP* (Upregulator of Cell Proliferation) was one such gene. *URGCP* is a cell-cycle regulator that drives proliferation via Wnt/ β -catenin signaling [25], and its reduction in the CNR-401 group is particularly noteworthy. In ALS, excessive proliferation of glial cells (astrocytes and microglia)

362 contributes to neuroinflammation and scarring [26]; therefore *URGCP* downregulation may help
363 curb this pathological gliosis. Indeed, suppression of *URGCP*/Wnt signaling has been shown to
364 dampen microglial activation and neuroinflammation in the spinal cord [27], suggesting that
365 CNR-401 creates a more quiescent glial environment.

366

367 Another key gene, *DAO* (*D-amino acid oxidase*), was also markedly downregulated, consistent
368 with neuroprotective mechanisms observed in ALS models. It has been demonstrated that *DAO*
369 inactivation in astrocytes reduces D-serine-mediated excitotoxicity, protecting motor neurons
370 [28]. However, *DAO* loss-of-function mutations in neurons have been linked to elevated
371 D-serine and neurodegeneration [29], highlighting context-dependent effects. CNR-401-induced
372 *DAO* suppression may reflect astrocyte-specific modulation, a complex feedback mechanism, or
373 a time-dependent effect – for example, if BMAA exposure had suppressed D-serine availability,
374 lowering *DAO* could be a compensatory attempt to restore normal neurotransmission.
375 Additionally, CNR-401's other actions (e.g. anti-glutamatergic or antioxidant effects) might
376 buffer the impact of increased D-serine in neurons. These findings underscore the intricacy of
377 excitotoxic pathways in ALS and suggest that while CNR-401 broadly mitigates BMAA toxicity,
378 certain aspects like D-serine metabolism require further investigation. Despite this complexity,
379 the functional mobility data showed that CNR-401 treatment rescues BMAA-induced motor
380 deficits, suggesting that any potential risks associated with altered D-serine metabolism are
381 indeed outweighed by the compound's overall neuroprotective effects in vivo.

382

383 *MXRA5* (*Matrix Remodeling Associated 5*) was already slightly downregulated in
384 BMAA-exposed zebrafish compared to control (supplementary material), but treatment with

385 CNR-401 caused a much greater suppression. *MXRA5* is involved in extracellular matrix (ECM)
386 remodeling and can facilitate MMP-9 activation, contributing to breakdown of the blood-spinal
387 cord barrier and tissue damage [30]. The pronounced suppression of this gene therefore suggests
388 that CNR-401 enhances a protective response. This suppression likely reflects modulation of the
389 well-established TGF- β 1 pathway, which controls *MXRA5* expression and ECM homeostasis
390 [31]. Our GO enrichment analysis strongly supports this hypothesis by identifying extracellular
391 matrix organization processes among the most significantly enriched terms in the CNR-401
392 group, indicating that therapeutic effects are largely mediated by the restoration of ECM stability.
393 Reduced *MXRA5* expression is part of this larger protective modulation of the ECM, leading to
394 reduced extracellular proteolysis and a more intact neurovascular unit, thereby protecting motor
395 neurons from inflammatory infiltration and toxic exposure.

396

397 Both *HLA-DQA1* and *HLA-DQA2*, major *histocompatibility complex (MHC) class II immune*
398 *genes*, were also among the top downregulated transcripts. Both proteins modulate immune
399 responses with the HLA-DQA1 protein forming a functional complex with HLA-DQB1 to create
400 an antigen-binding heterodimer that displays foreign peptides derived from extracellular proteins
401 to the immune system [32], and HLA-DQA2 modulating immune responses through
402 non-classical pathways [33]. In the central nervous system, MHC class II molecules like
403 HLA-DQ are typically absent under healthy conditions but become upregulated on microglia
404 during neuroinflammatory processes, making them key markers of activated immune cells in the
405 brain [34]. Lower *HLA-DQ* expression therefore suggests suppression of neuroimmune
406 activation in treated zebrafish. This aligns with CNR-401's general anti-inflammatory signature,
407 as reduced *MHC II* expression limits the activation of CD4⁺ T cells and the transition of

microglia from a resting to a reactive, pro-inflammatory state in the CNS. Along with *URGCP* and *MXRA5*, the downregulation of *HLA-DQA1* and *HLA-DQA2* reinforces the notion that CNR-401 dampens processes that contribute to neuroinflammation and the neurodegenerative environment.

In contrast to the many downregulated genes, *UNG* (*Uracil-DNA glycosylase*) stood out as one of the few upregulated genes. *UNG* is a DNA repair enzyme involved base-excision repair that removes uracil from DNA, thereby correcting cytosine deamination and other oxidative DNA damage [35]. Its upregulation by CNR-401 suggests an enhanced capacity for genomic maintenance in response to BMAA toxicity and associated oxidative stress, which inflicts DNA damage in neurons [36]. Given that accumulated DNA damage in motor neurons is a known contributor to ALS progression [37], this pro-repair response would be neuroprotective.

Collectively, the top DEGs indicate that CNR-401 treatment modulates key players in detoxification and steroid metabolism (*UGT* genes), cell proliferation/inflammation (*URGCP*, *HLA-DQA1*, *HLA-DQA2*), excitotoxic signaling (*DAO*), matrix integrity (*MXRA5*), and genomic stability (*UNG*). Such changes are broadly consistent with a therapeutic shift toward neuroprotection – reducing toxic exposures, calming injurious immune activity, preserving structural support, and bolstering repair mechanisms.

Enriched Biological Processes: ECM Organization and Inflammation

The differential expression patterns translate into clear enrichment of biological processes and pathways relevant to ALS. The Gene Ontology enrichment analysis reveals a compelling convergence of two fundamental pathological processes that serve as immediate contributors to

431 motor neuron degeneration in ALS: inflammatory response and extracellular matrix (ECM)
432 organization. These processes represent the most direct mechanisms by which BMAA initiates
433 motor neuron injury and how CNR-401 subsequently counteracts these damaging processes.
434

435 The enrichment of ECM organization terms “extracellular matrix organization”, “external
436 encapsulating structure organization”, and “extracellular structure organization” in our analysis
437 directly reflects the mechanistic foundation of ALS pathology described in the literature.
438 Research of the related neurodegenerative disease multiple sclerosis (MS) has demonstrated that
439 many ECM components elevated during tissue damage are inherently pro-inflammatory and
440 enhance inflammatory processes [38]. Within the context of ALS, breakdown of the extracellular
441 matrix represents a critical initiating event that triggers a cascade of inflammatory responses
442 leading to motor neuron degeneration [39]. In healthy neural tissue, stable ECM and basement
443 membranes provide structural support that maintains microglia in a ramified, quiescent state.
444 However, when ECM breakdown occurs, microglia become mobile and reactive, further
445 degrading the already compromised extracellular matrix and establishing a chronic
446 para-inflammatory state in the motor tracts [39]. This creates a self-perpetuating cycle where
447 ECM damage promotes inflammation, which in turn exacerbates ECM degradation.
448

449 This prominent enrichment of inflammatory response processes was captured in our Gene
450 Ontology analysis, which showed "inflammatory response", "cytokine-mediated signaling
451 pathway", and "cellular response to cytokine stimulus" among the most significant terms. This
452 provides compelling evidence for neuroinflammation's central role in ALS pathogenesis. The
453 convergence of inflammatory processes aligns with extensive research that demonstrates

454 neuroinflammation as a fundamental mechanism that drives motor neuron injury rather than
455 merely a secondary consequence of neurodegeneration. Motor neuron degeneration in ALS
456 operates through non-cell autonomous mechanisms, where motor neuron death results not solely
457 from intrinsic cellular defects but from active contributions of surrounding glial cells. This
458 concept has been definitively established through chimeric mouse studies demonstrating that
459 wild-type neurons acquire ALS phenotypes when surrounded by glial cells expressing mutant
460 proteins, confirming that the cellular environment directly determines motor neuron survival
461 [40]. Neuroinflammation in ALS is characterized by coordinated activation of multiple immune
462 and glial populations that directly contribute to motor neuron degeneration. This inflammatory
463 milieu includes infiltration of peripheral lymphocytes and macrophages, widespread activation of
464 resident microglia and reactive astrocytes, and engagement of complement cascades that
465 collectively contribute to motor neuron death through the coordinated release of
466 pro-inflammatory cytokines, reactive oxygen species, and other neurotoxic factors [41]. The
467 neuroinflammatory response transforms the normally supportive glial environment into one that
468 actively promotes motor neuron degeneration, establishing neuroinflammation as a primary
469 therapeutic target for disease-modifying interventions in ALS [41].

470

471 The observation that inflammatory response and ECM organization are enriched processes in
472 both BMAA toxicity and CNR-401 treatment provides compelling evidence for a coherent
473 pathological and therapeutic narrative. This shared enrichment demonstrates that these biological
474 processes constitute the core molecular battleground where neurodegeneration occurs and where
475 therapeutic intervention must be focused. The initial exposure to BMAA triggers ECM
476 degradation and neuroinflammation that directly damages motor neurons. Then, subsequent

477 CNR-401 treatment appears to target these same fundamental pathways, suggesting that this
478 therapeutic specifically addresses the root mechanisms of BMAA-induced motor neuron injury.
479 While BMAA disrupts these systems through toxic mechanisms, CNR-401's multi-target
480 botanical approach modulates the same pathways through protective mechanisms. The
481 therapeutic enrichment of these pathways likely represents CNR-401's dual capacity to stabilize
482 ECM integrity while simultaneously modulating inflammatory responses to create a more
483 permissive environment for motor neuron survival and potential regeneration. This convergence
484 of pathological initiation and therapeutic intervention on identical biological processes
485 underscores the mechanistic precision of CNR-401's therapeutic approach and validates the
486 fundamental importance of ECM organization and inflammatory response as direct contributors
487 to motor neuron injury in ALS.

488

489 Enriched Biological Processes: Calcium Signaling and Steroid Metabolism

490 The Gene Ontology enrichment of calcium-mediated signaling and steroid metabolic processes
491 represents a secondary tier of regulatory mechanisms that modulate cellular stress responses and
492 provide neuroprotective functions in ALS pathogenesis. These pathways operate as crucial
493 modulators of cellular homeostasis, influencing both the susceptibility to neurodegeneration and
494 the capacity for neuroprotective responses following initial ECM and inflammatory damage.

495

496 Motor neurons in ALS exhibit fundamental vulnerabilities in calcium homeostasis that
497 predispose them to calcium-mediated toxicity. These neurons express high levels of
498 calcium-permeable AMPA receptors while maintaining low calcium buffering capacity, creating
499 susceptibility to intracellular calcium overload [42]. This calcium dysregulation operates through

500 a "toxic shift" [43] within the endoplasmic reticulum-mitochondria calcium cycle, representing a
501 key mechanism in motor neuron degeneration. The chronic excitotoxicity mediated by AMPA
502 receptors initiates a self-perpetuating process of intracellular calcium dysregulation with
503 consecutive endoplasmic reticulum calcium depletion and mitochondrial calcium overload,
504 ultimately spiraling toward catastrophic levels of cytosolic calcium [43, 44]. The enrichment of
505 "calcium-mediated signaling" in the CNR-401 versus BMAA comparison provides molecular
506 evidence for CNR-401's potential therapeutic effects through modulation of these fundamental
507 cellular stress response mechanisms that are compromised in ALS pathology.

508

509 The emergence of steroid metabolic process and specifically estrogen metabolic process in GO
510 enrichment correlates with the downregulation of *UGT (UDP-glucuronosyltransferase)* enzymes
511 that conjugate steroid hormones, and reflects CNR-401's anti-catabolic influence on critical
512 endogenous neuroprotective mechanisms. The central nervous system possesses capacity for
513 local steroid synthesis, producing progesterone, testosterone, and estradiol that provide
514 neuroprotective effects independent of peripheral hormonal sources [45, 46]. For example, one
515 such neurosteroid is allopregnanolone, a progesterone metabolite that demonstrates
516 neuroprotective properties through mechanisms including GABA-A receptor modulation,
517 anti-apoptotic signaling, and neuroinflammation regulation [46, 47]. Estrogen metabolism
518 pathways are particularly relevant to ALS, demonstrating anti-apoptotic mechanisms, calcium
519 homeostasis regulation, microglial modulation, anti-inflammation, and antioxidant activities [48,
520 49]. Specific examples include 17 β -estradiol which demonstrates direct protective effects on
521 spinal motor neurons through anti-apoptotic and anti-inflammatory actions on glial cells [50],
522 and estrogen signaling mediated by G-protein-coupled receptor 30 (GPR30) which demonstrates

523 similar anti-apoptotic effects on motor neurons [51]. The downregulation of steroid-conjugating
524 enzymes by CNR-401 suggests a sophisticated therapeutic strategy that enhances the
525 bioavailability and duration of action of endogenous neuroprotective steroids. This mechanism
526 provides a molecular explanation for how botanical therapeutics can enhance natural
527 neuroprotective responses without requiring exogenous hormone supplementation.

528

529 The enrichment of both calcium and steroid processes suggests that CNR-401 targets these
530 interconnected regulatory systems to provide comprehensive neuroprotective effects. By
531 modulating calcium homeostasis while potentially enhancing neurosteroid bioavailability,
532 CNR-401 appears to restore cellular capacity for adaptive responses to pathological stimuli while
533 enhancing endogenous neuroprotective mechanisms that are compromised in ALS progression.
534 These enriched terms represent sophisticated therapeutic targets that address multiple levels of
535 cellular dysfunction, explaining how botanical therapeutics can simultaneously target
536 excitotoxicity, neuroinflammation, and cellular stress responses.

537 KEGG Pathway Enrichment Reinforces GO Biological Process and Gene Expression 538 Findings

539 KEGG pathway enrichment analysis reinforces these GO biological process findings. After
540 CNR-401 treatment, “cytokine–cytokine receptor interaction” was the top pathway, directly
541 supporting the GO enrichment of inflammatory response processes and confirming that
542 CNR-401’s therapeutic mechanism involves comprehensive modulation of immune signaling
543 networks that drive neuroinflammation in ALS pathogenesis. This pathway enrichment indicates

544 that CNR-401 affects not only individual cytokine expression but the broader intercellular
545 communication systems that perpetuate chronic neuroinflammatory states.

546

547 Furthermore, the enrichment of the "steroid hormone biosynthesis" and "ovarian steroidogenesis"
548 pathways directly reinforces the previous interpretation that CNR-401 enhances the
549 bioavailability and persistence of endogenous neuroprotective steroids. These KEGG pathways
550 capture the broader metabolic context in which UGT downregulation operates, highlighting that
551 the transcriptomic impact of CNR-401 extends beyond isolated gene effects to coordinated
552 modulation of entire steroidogenic networks. This systemic enrichment supports the conclusion
553 that CNR-401 not only reduces steroid catabolism via *UGT* suppression but also potentially
554 upregulates or stabilizes upstream biosynthetic processes, thereby amplifying the pool of
555 neuroactive steroids available to exert anti-inflammatory, anti-apoptotic, and neuroprotective
556 functions relevant to ALS pathophysiology.

557

558 Appearance of "ascorbate and aldarate metabolism" and "retinol metabolism" among enriched
559 pathways provides mechanistic insight into CNR-401's antioxidant capabilities, complementing
560 the GO enrichment of calcium signaling and steroid metabolic processes, and the marked
561 downregulation of *DAO*. Ascorbate (vitamin C) is a major direct antioxidant and free radical
562 scavenger in the central nervous system [52, 53], playing a critical role in neutralizing reactive
563 oxygen species and protecting neurons from the oxidative stress in ALS pathogenesis. Similarly,
564 retinol metabolism enrichment indicates modulation of vitamin A, an indirect antioxidant [54].
565 Both vitamins A and C have been highlighted as natural antioxidants with the potential to
566 suppress neural oxidative damage and slow neurodegeneration in ALS [55].

567

568 Two metabolic pathways—"pentose and glucuronate interconversions" and "metabolism of
569 xenobiotics by cytochrome P450"—directly relate to the detoxification effect and the
570 coordinated downregulation of UGT enzymes discussed previously. The pentose and glucuronate
571 interconversions pathway represents a fundamental phase II detoxification mechanism [56] and
572 the central biochemical process within this pathway, glucuronidation, is primarily catalyzed by
573 UDP-glucuronosyltransferase (UGT) enzymes [57]. This evidence thus demonstrates that
574 CNR-401 fundamentally alters cellular detoxification capacity by targeting both the enzymatic
575 machinery (UGT downregulation) and the broader metabolic framework (pathway enrichment)
576 of Phase II conjugation processes. Transcriptomic analyses across multiple species have
577 consistently identified metabolism of xenobiotics by cytochrome P450 as a major Phase I
578 pathway intersecting with the broader detoxification processes [58, 59]. The enrichments of these
579 two pathways thus provide systems-level evidence that CNR-401 reduces the metabolic burden
580 of toxin processing, with cytochrome P450 enzymes catalyzing Phase I oxidative metabolism
581 and glucuronidation serving as a critical Phase II conjugation step that enhances the water
582 solubility and excretion of metabolites [59].

583

584 Novel Pathway Identified by KEGG and Comparison to Control

585 Beyond validating GO enrichment findings, KEGG analysis revealed additional therapeutic
586 mechanisms not captured in biological process enrichment. The enrichment of the PI3K-Akt
587 signaling pathway represents a critical finding, as this pathway serves as a central regulator of
588 neuronal survival and has been extensively implicated in neuroprotective mechanisms across
589 numerous neurodegenerative diseases [60]. The PI3K-Akt pathway plays a pivotal role in

neuroprotection by promoting cell survival through stimulation of cell proliferation and inhibition of apoptosis, and modulation of cellular stress responses including oxidative stress and inflammatory signaling [61]. In ALS pathogenesis specifically, studies using SOD1-G93A ALS mouse models have shown that treatments enhancing PI3K-Akt signaling reduce disease progression by improving motor function and reducing pathological changes [62]. This enrichment therefore suggests that CNR-401's neuroprotective effects partially operate through activation of survival signaling cascades that counteract BMAA-induced oxidative stress, inflammatory responses, and apoptotic cell death.

For contextual comparison, the BMAA versus control enrichment included terms such as JAK-STAT signaling pathway, P13K-AKT signaling pathway, cytokine-cytokine receptor interaction, and ECM-receptor interaction, confirming that BMAA exposure induces inflammatory and destructive responses characteristic of neurodegenerative injury. The therapeutic enrichment patterns in CNR-401 treatment directly counteract these pathological processes, demonstrating mechanistic specificity in addressing BMAA-induced molecular perturbations.

The convergence of KEGG pathway and GO enrichment analyses establishes that CNR-401's neuroprotective mechanism operates through coordinated modulation of inflammation (cytokine signaling), steroid bioavailability (steroid biosynthesis), antioxidant defense (vitamin metabolism), detoxification (glucuronate and xenobiotic metabolism), and cellular survival (PI3K-Akt)—precisely targeting pathological processes that drive ALS progression.

612 Comparing to Edaravone and Cannflavin A: Distinct Mechanistic Signatures

613 The differences between CNR-401's transcriptomic effects and those of Edaravone or Cannflavin
614 A shed light on their divergent mechanisms of action. Edaravone, an FDA-approved ALS drug,
615 is a well-known free radical scavenger that provides neuroprotection by reducing oxidative stress
616 and delaying motor neuron deterioration [63, 1]. Its mechanism operates through direct chemical
617 neutralization of reactive oxygen species, making it fundamentally a non-genomic intervention
618 rather than a transcriptional modulator. In our study, Edaravone treatment after BMAA exposure
619 led to relatively few DEGs (only 359, with the vast majority being downregulations). However,
620 this limited transcriptomic footprint does not align with precise targeting of oxidative stress
621 pathways as the literature claims. Our pathway enrichment analysis revealed no significant
622 enrichment of oxidative stress-related pathways and instead revealed metabolic pathways
623 including steroid biosynthesis, lipid metabolism, and alcohol metabolism, alongside cell
624 signaling pathways such as PI3K-Akt, VEGF, and JAK-STAT signaling (supplementary
625 material). This limited yet scattered transcriptomic footprint may explain why Edaravone
626 provides only modest clinical benefits despite FDA approval [1]; The scattered pattern suggests
627 that edaravone lacks the focused potency needed for robust therapeutic intervention. CNR-401,
628 by contrast, alters hundreds of genes across diverse pathways in a more coordinated fashion,
629 suggesting a more comprehensive and effective therapeutic mechanism that addresses multiple
630 aspects of neurodegeneration simultaneously rather than superficially touching many pathways.
631

632 Cannflavin A, a flavonoid from *Cannabis sativa*, was included as another comparator. In our
633 zebrafish ALS model, pure Cannflavin A treatment at 2 μ M after BMAA exposure resulted in
634 130 differentially expressed genes (DEGs), with a predominant upregulation pattern (91 up vs 39

635 down, supplementary material). Remarkably, the enrichment analysis revealed that Cannflavin A
636 did not significantly enrich inflammatory response pathways, despite its well-established
637 anti-inflammatory properties [13, 14]. Our enrichments showed that it instead activated
638 metabolic, antioxidant, and steroidogenic pathways (supplementary material). This apparent
639 discrepancy can be explained by critical pharmacological and mechanistic factors that highlight
640 the complexity of natural product bioactivity translation from *in vitro* to *in vivo* systems. While
641 our 2 μ M concentration theoretically exceeds the reported cell-free (*in-vitro*) IC50 values for
642 these enzymes (mPGES-1 inhibition at 1.8 μ M and 5-LOX inhibition at 0.9 μ M) [13], *in vivo*
643 bioavailability factors, which are known to significantly reduce effective tissue concentrations
644 [64], may have prevented proper inhibition. Critically, Cannflavin A treatment still improved
645 locomotor function in BMAA-exposed zebrafish, demonstrating functional neuroprotective
646 efficacy despite the absence of classical inflammatory pathway modulation. Regardless of
647 through which mechanisms, the scope of changes with Cannflavin A was still limited compared
648 to CNR-401. Notably, Cannflavin A did not trigger the concerted downregulation of UGT
649 enzymes that CNR-401 did, nor did it upregulate DNA repair enzymes or modulate the
650 P13K/Akt pathway, suggesting that these effects are unique to CNR-401. These distinct
651 transcriptomic signatures allow us to directly compare how CNR-401's mechanism differs from
652 these known therapies. Edaravone's signature is one of a scattered modulator with limited effects
653 across diverse pathways and Cannflavin A's signature (at this concentration) was that of a
654 focused modulator of metabolic and steroidogenic functions. CNR-401, however, produced a
655 hybrid signature by combining these elements plus anti-inflammation, regulation of calcium
656 overload, cellular survival through PI3K-Akt signaling, and even DNA-reparative effects.

657 CNR-401 Has Superior Anti-inflammatory Activity

658 When CNR-401 was tested at a concentration of 2 μ M, the compound demonstrated robust
659 anti-inflammatory gene expression signatures, including significant downregulation of cytokine
660 signaling pathways and *MHC-II* genes (*HLA-DQA1/DQA2*). This anti-inflammatory
661 transcriptomic response is particularly noteworthy given that pure Cannflavin A, when tested at
662 the same 2 μ M concentration, failed to enrich canonical inflammatory-response pathways despite
663 its well-established anti-inflammatory properties. The complex botanical composition of
664 CNR-401 may contribute to its broader and more pronounced modulation of inflammatory gene
665 networks through the additive effects of its various components, and potentially through
666 synergistic interactions between them. Although the current study is not designed to assess
667 synergy, such effects have been proposed between cannabinoids and terpenes in various other
668 therapeutic areas [16]. What can be said about these results is that CNR-401 achieves therapeutic
669 anti-inflammatory effects at total concentrations where Cannflavin A would be insufficient,
670 supporting a sophisticated combinatorial strategy that maximizes therapeutic benefit while
671 potentially minimizing the concentration requirements of any single active ingredient.

672 Future Research Directions

673 Interpretation of transcriptomic data requires careful consideration as a more extensive change in
674 gene expression (as seen with CNR-401) does not automatically equate to superior clinical
675 efficacy; some changes could be neutral or even maladaptive. The observed *DAO*
676 downregulation in CNR-401-treated subjects exemplifies this complexity. While functional
677 mobility assessments demonstrated that CNR-401 treatment rescues rather than exacerbates
678 BMAA-induced motor deficits, examples like *DAO* emphasize the critical importance of

679 integrating transcriptomic and functional analyses to accurately evaluate the net therapeutic
680 impact of candidate interventions on disease-relevant outcomes. Although CNR-401
681 demonstrated a positive net therapeutic effect in this study, the extensive genetic modifications
682 warrant detailed mechanistic investigation to establish clinical relevance for each significant
683 genetic change.

684

685 Edaravone's limited transcriptomic signature yet comparable functional outcome seems to
686 suggest that focused molecular interventions can achieve meaningful therapeutic outcomes with
687 minimal transcriptomic disruption. Despite inducing far fewer differentially expressed genes than
688 CNR-401 (359 versus 1,576 DEGs), Edaravone produced comparable functional rescue in our
689 mobility assessments. This indicates that strategic modulation of key pathways may be as
690 effective as broad transcriptomic reprogramming. However, this pattern may simply reflect the
691 limitations of our BMAA-induced ALS model in revealing the full therapeutic potential of
692 CNR-401's multi-pathway modulation. To determine whether CNR-401's extensive genetic
693 changes confer additional therapeutic advantages compared to Edaravone, both compounds
694 should be tested across diverse experimental conditions that challenge different aspects of ALS
695 pathophysiology, such as chronic neurodegenerative models and genetic ALS variants. The
696 potential value of CNR-401's extensive transcriptomic modulation may only become apparent
697 under these specific disease conditions or patient contexts not captured by our current model,
698 emphasizing the critical importance of comprehensive preclinical testing to fully evaluate its
699 therapeutic potential.

700

701 Given that the transcript-level changes observed with CNR-401 are now matched by robust
702 functional neuroprotection (the significant rescue of locomotor activity in BMAA-exposed
703 zebrafish), CNR-401 emerges as a promising next-generation multi-target therapeutic candidate
704 for ALS. The convergence of molecular and behavioral efficacy strengthens the rationale for
705 further preclinical development and positions CNR-401 as a lead compound capable of
706 addressing the complex, multifactorial pathology of ALS. Future studies should validate these
707 multi-target genetic interpretations by measuring corresponding biochemical changes, such as
708 neurosteroid levels or oxidative damage markers, in CNR-401-treated subjects. Additionally,
709 comparing CNR-401's transcriptomic signature to those from other ALS models or
710 patient-derived cells could further clarify how this compound fits into the landscape of ALS
711 therapeutics. Nonetheless, the current results are encouraging: they suggest that CNR-401
712 intervenes in multiple downstream consequences of ALS pathology, potentially offering a
713 broader therapeutic benefit than existing single-mechanism drugs.

714

715 CONCLUSIONS

716 Summary of CNR-401's Neuroprotective Mechanisms

717 Our transcriptomic analysis reveals that CNR-401 addresses multiple pathological processes
718 underlying ALS through a coordinated molecular response. The compound demonstrates
719 multi-targeted neuroprotection by simultaneously modulating eight key mechanisms:

720

721 **Neuroinflammation Suppression and ECM Stabilization:** CNR-401 downregulates key
722 inflammatory mediators including URGCP and HLA-DQA1/DQA2, while matrix remodeling

723 gene MXRA5 shows pronounced suppression. GO enrichment reveals terms related to
724 extracellular matrix organization and inflammatory/immune response, while KEGG pathway
725 analysis identifies "cytokine-cytokine receptor interaction" and "ECM-receptor interaction" as
726 significantly enriched, confirming comprehensive modulation of immune signaling networks and
727 ECM stabilization.

728

729 **Detoxification Pathway Modulation:** Coordinated downregulation of 19 human UGT orthologs
730 represents the most striking transcriptomic signature and KEGG pathways "pentose and
731 glucuronate interconversions" and "metabolism of xenobiotics by cytochrome P450" are
732 significantly enriched, indicating fundamental alterations in Phase I and Phase II detoxification
733 processes.

734

735 **Neurosteroid Biosynthesis Enhancement:** The coordinated UGT downregulation directly
736 impacts neurosteroid bioavailability through reduced conjugation, supported by GO and KEGG
737 enrichments of steroid and estrogen metabolic processes. This enhances production and
738 persistence of neuroactive steroids with anti-inflammatory and neuroprotective properties.

739

740 **Excitotoxicity Modulation:** CNR-401's downregulation of DAO likely reflects
741 astrocyte-specific effects or compensatory mechanisms that contribute to CNR-401's
742 comprehensive approach to addressing excitotoxic stress in ALS pathogenesis.

743

744 **Antioxidant Defense Enhancement:** KEGG pathway enrichment reveals "ascorbate and
745 aldarate metabolism" and "retinol metabolism" as significantly enriched pathways, indicating
746 enhanced antioxidant capacity through vitamin C and vitamin A metabolism.

747 - Regulation of Calcium Overload: GO enrichment of "calcium-mediated signaling" indicates
748 that CNR-401 modulates calcium buffering to prevent calcium overload. This regulation
749 addresses the fundamental "toxic shift" in calcium homeostasis that characterizes motor neuron
750 death.

751

752 **DNA Repair Enhancement:** UNG (Uracil-DNA glycosylase) upregulation indicates enhanced
753 base-excision repair capacity, representing a mechanism not addressed by current ALS therapies.
754 This suggests improved genomic maintenance capacity against oxidative DNA damage.

755

756 **Cellular Survival Through PI3K-Akt Signaling:** KEGG enrichment of "PI3K-Akt signaling
757 pathway" represents a critical finding, as this pathway serves as a central regulator of neuronal
758 survival and neuroprotective mechanisms across neurodegenerative diseases. This suggests
759 CNR-401's effects operate through activation of survival signaling cascades that counteract
760 oxidative stress and apoptotic cell death.

761

762 This comprehensive molecular analysis demonstrates that CNR-401's neuroprotective efficacy
763 stems from coordinated multi-pathway modulation rather than single-target effects. Critically,
764 these transcriptomic changes translate into functional neuroprotection, as demonstrated by
765 significant rescue of locomotor deficits in BMAA-exposed zebrafish, confirming that molecular
766 modulation preserves motor outcomes in this ALS model.

767

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771

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775

776 **AUTHOR CONTRIBUTIONS**

777 • **K.B., E.S., and A.G.** conceived and designed the experiments.

778 • **K.B. and E.S.** carried out the experiments.

779 • **J.C.** provided scientific advisory throughout all steps.

780 • **I.B. and G.D.** analyzed the data.

781 • **I.B.** wrote the manuscript.

782 • All authors read and approved the final manuscript.

783

784 **DATA AND CODE AVAILABILITY**

785 All data generated and analyzed during this study are provided as in the Supplementary Materials

786 which can be found at

787 [https://figshare.com/projects/Transcriptomic_Analysis_of_Neuroprotective_Therapies_in_a_Zeb](https://figshare.com/projects/Transcriptomic_Analysis_of_Neuroprotective_Therapies_in_a_Zebrafish_Model_of_ALS/251624)

788 rafish_Model_of_ALS/251624. This includes the product toxicity assay results, original

789 normalized RNA-seq data, and the complete set of downstream differential expression data for

all 20 experimental comparisons. Additional figures such as volcano plots, MA plots, heatmaps, and enrichment barplots are also available in the supplementary materials.

All code for the pipeline used for analysis (the DEG Pipeline Assistant) is available at <https://github.com/Shaan7071/DEG-pipeline-assistant>. Further information or materials can be requested from the corresponding author.

CONFLICTS OF INTEREST

This research was funded by Canurta Therapeutics, to which the authors belong.

ETHICAL APPROVAL

No formal animal ethics approval was required for this study, as all zebrafish experiments were performed in compliance with relevant institutional and national guidelines for the care and use of laboratory animals.

CONSENT TO PARTICIPATE

Ethics and consent were not required for the performed study.

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