Cellular and molecular profiles of anterior nervous system regeneration in *Diopatra claparedii* Grube, 1878 (Annelida, Polychaeta)

Mohd Ulul Ilmie Ahmad Nazria,b, Mohd Hafizi Mahmudc, Basri Saidic, Mohd Noor Mat Isa d, Zolkapli Ehsake, Othman Rossf, Izwandy Idrisg, Wan Iryani Wan Ismaila,b,*

a Cell Signalling and Biotechnology Research Group (CeSBTech), Faculty of Science and Marine Environment, Universiti Malaysia Terengganu, 21030, Kuala Nerus, Terengganu, Malaysia
b Biological Security and Sustainability Research Group (BIOSSES), Faculty of Science and Marine Environment, Universiti Malaysia Terengganu, 21030, Kuala Nerus, Terengganu, Malaysia
c Centre of Medical Imaging, Faculty of Health Sciences, Universiti Teknologi MARA Selangor, Puncak Alam Campus, 42300, Bandar Puncak Alam, Selangor, Malaysia
d Malaysia Genome Institute, Jalan Bangi, 43000, Kajang, Selangor, Malaysia
e Imaging Centre, Faculty of Pharmacy, Universiti Teknologi MARA Selangor, Puncak Alam Campus, 42300, Bandar Puncak Alam, Selangor, Malaysia
f Faculty of Science and Marine Environment, Universiti Malaysia Terengganu, 21030, Kuala Nerus, Terengganu, Malaysia
g South China Sea Repository & Reference Centre, Institute of Oceanography & Environment, Universiti Malaysia Terengganu, 21030, Kuala Nerus, Terengganu, Malaysia

ARTICLE INFO

Keywords:
Annelida
Nervous system
Early regeneration
Polychaete
Ultrastructure
Transcriptome
Neurodegenerative diseases
Micro CT scan
Ultramicrotome
Bioinformatics

ABSTRACT

The polychaete *Diopatra claparedii* Grube, 1878 is among those organisms successfully carrying out full body regeneration, including the whole nervous system. Thus, *D. claparedii* potentially can be regarded for the nervous system regeneration (NSR) study. However, data on the property of its nervous system and the NSR profile are still lacking. In this study, we investigated the morphology of *D. claparedii* anterior nervous system (ANS) and examined the cellular and molecular profiles on its early anterior NSR. The nervous system of *D. claparedii* consists of a symmetry brain with nerves branching off, circumpharyngeal connectives that connect the brain and nerve cord as well as obvious segmental ganglia. Moreover, we identified changes in the cellular condition of the ganglionic cells in the regenerating tissue, such as the accumulation of lysosomes and lipofuscins, elongated mitochondria and multiple nucleoli. Furthermore, mRNA of tissues at two regenerating stages, as well as intact tissue (non-regenerating), were sequenced with Illumina sequencer. We identified from these tissues 37,248 sequences, 18 differential expressed proteins of which upregulated were involved in NSR with noelin-like isoform X2 turned up to be the highest being expressed. Our results highlight the cellular and molecular changes during early phase of NSR, thus providing essential insights on regeneration within Annelida and understanding the neurodegenerative diseases.

1. Introduction

Nervous system regeneration (NSR), or also known as neuroregeneration, is defined as restoration and regrowth of the lost neuron or nerve tissue to regain normal function. In invertebrates, such as echinoderms, the regeneration capacity is remarkable as they can regenerate both internal and external parts, including the nervous system (Ortega and Olivares-Bañuelos, 2020). Neurogenesis is a process by which a new neuron is produced from a stem/precursor cell during the developmental process from juvenile to adult. However, in some adult insects, it was found that persistent neurogenesis occurs in the mushroom bodies which are known to function in learning and memory (Cayre et al., 2002). Furthermore, a study conducted on an adult serpulid polychaete has demonstrated the outgrowth of a cerebral ganglion as the first feature established during neurogenesis (Brinkmann and Wanninger, 2009). On the other hand, the second process, neuroplasticity, is an adaptation of the brain towards environment enhancement in terms of structure and function (Enciu et al., 2011). In invertebrates, especially mollusks, and insects, neuroplasticity has been reported to occur for axon sprouting and synapse formation (Pysz, 2013). Thus, the cause of neurodegeneration, on the contrary, might be due to the absence of the neuroplasticity process.

The capacity for neuroregeneration varies widely among taxa. Invertebrates are known to regenerate the entire body while...
phylogenetically primitive vertebrates such as lizards and zebrafish selectively regenerate certain parts. Lizards such as geckos and skinks, for example, regenerate the losing tail once the tail drops. However, the regenerated tail remains different because the original spinal cord is replaced by an epithelial tube (Alibardi, 2009). Meanwhile, the zebrafish, *Danio rerio* is widely used as an animal model to study neurodegenerative disease. The zebrafish is able to repair lesions in the brain, regenerate tissue, and neurons in the spinal cord, brain, and retina, but only if the residual glial cells remain (Gemberling et al., 2013; Nazri et al., 2019). However, the regeneration does not occur de novo. Thus, several organisms, mostly invertebrates, maintain the ability to regenerate certain parts for survivability. Moreover, some invertebrate species are capable of regenerating entirely to become new individuals, for example, flatworm (Platyhelminthes) and marine worms from class Polychaeta (phylum Annelida). In polychaetes, NSR begins with the outgrowth of nerve fibre from the nerve cord of the original body that later formed the brain and other nerves (Weidhase et al., 2015).

Polychaetes represent almost 65% of all annelid species (Anderson, 2013) making them an incredibly diverse and abundant marine invertebrate group. The class was erected by Grube (1850) and, at present, consists of 85 families. Most of them are benthic, and some are pelagic, which can be found in marine, brackish water; a small number of species live in freshwater and even less, terrestrial (Thorp and Rogers, 2014). Polychaetes are separated into two large orders, Errantia and Sedentaria based on their morphology and behaviour. Errant polychaetes are highly mobile and free-living while sedentary polychaetes are typically tubicolous. Their adult size varies from less than 1 mm up to 6 m. Their colours are often vibrant and iridescent. They feed on smaller prey or by trapping food particles by ciliary action. Polychaetes reproduce sexually with separate sexes and some asexually with a high number of variabilities. Fertilisation mainly occurs externally in the water and results in either free-swimming larvae or direct development (Rouse and Pleijel, 2013) making them an incredibly diverse and abundant marine invertebrate group.

2. Materials & methods

2.1. Animals

The polychaetes were collected in the intertidal mudflat area in Morib Beach, Banting (Figure 1), west coast of Peninsular Malaysia (2.7495° N, 101.4426° E). The worms were removed by digging the sandy mud sediment to obtain the longest possible tubes. Each collected tube was checked to make sure the worm is present. The worms are susceptible to sediment vibration caused by human footprint and will burrow deeper into the sediment. Hence, the minimum length of *D. claparedii* that can be used for the experiment is 5 cm when relaxing. For the laboratory acclimatisation, all collected *D. claparedii* were maintained in the aquaria containing sediment from the sampling site for two weeks, at 28 °C and 23–25 ppt salinity of artificial seawater (Aquaforest; Brzesko, Poland). The worms were fed with commercial sinking fish pallet *ad libitum*. During the habituation period, the polychaetes were allowed to regenerate the posterior section themselves since they were accidentally amputated during collection. In each experimental procedure, the polychaetes were immobilised with 4% MgCl2.6H2O.

2.2. Cresyl violet staining

Intact polychaetes were fixed in 10% formalin mixed with artificial seawater overnight (Mackie, 1994). Then, the fixed specimens were dehydrated in ascending alcohol (70–100%) to remove all traces of water, followed by clearing in xylene. The anterior section (prostomium and peristomium) and all external appendages were then removed. The anterior part was then embedded in paraffin wax prior to sectioning. The paraffin block was sectioned to 5 μm of thickness using a microtome and then proceed to the standard histological slide preparation before staining. The slides were then immersed in 0.5% cresyl violet, which stained the Nissl bodies in the neurons for 2 min and washed again with distilled water followed by dehydration by 70% ethanol and twice absolute ethanol for 2 min, respectively and finally with a 5 min wash in

![Figure 1. Map of the study with the location of sampling site in Morib Beach, Banting, western Peninsular Malaysia.](image-url)
xylene. The sections were covered up with a coverslip and viewed under a light compound microscope.

2.3. Immunohistochemistry combining with confocal laser scanning microscopy

The anterior part of intact polychaetes was fixed in 4% paraformaldehyde for 12 h, before overnight incubation in 15% sucrose solution (to sustain the tissue shape). The specimen was then mounted on OCT medium before sectioning by cryostat Leica CM1950 (Leica Biosystems, Wetzlar, Germany). The specimen was sectioned 8 μm thickness horizontally in dorsal position for the brain and ventral position for ventral nerve cord. The sections were then transferred onto a superfrrost glass slide (Thermo Scientific, Waltham, MA, USA). The staining procedure was conducted according to the protocol from Cell Signaling Technology with several modifications. The sections were fixed in an ice-cold (-80 °C) methanol for 10 min, followed by two washes with PBS for 5 min in each wash. Next, the sections were incubated in 3% H2O2 diluted in methanol for 10 min at room temperature (RT) before two washes by PBS. The samples were then blocked for 1 h at RT by PBST-NGS (0.1 M PBS, goat serum (Sigma), triton-X with the ratio of 200:4:1). The samples were then incubated with primary antibody (monoclonal anti-acetylated α-tubulin by Santa Cruz, Dallas, TX, USA; dilution 1:250 in PBST-NGS) in a humid dark box at 4 °C. The sections were then washed twice with PBS for 5 min each and incubated with secondary antibody (polyclonal mouse IgG conjugated DyLight488 by GeneTex, Irvine, CA, USA; dilution 1:500) for 30 min at RT in the same dark box. Finally, the samples were washed before being mounted with an antifading mounting medium and covered with a coverslip. The slides containing sections were viewed with a Leica TCS SPE confocal microscope (Leica Biosystems) at 488 nm wavelength.

2.4. Micro-computed tomography scanning (Micro-CT)

Intact polychaetes were dehydrated in a series of ascending ethanol for 45 min each (90% twice, 96% twice) before they were dried with hexamethyldisilazane (HMDS) for 2 h. The specimens were left overnight prior to tomography scanning. Image of the specimen was acquired by using SkyScan 1176 microtomography (Bruker micro-CT; Kontich, Belgium) in medical imaging laboratory at Faculty of Health Sciences, UiTM Puncak Alam, Selangor. Specimens were scanned at 9 μm resolution. A 50 kV voltage with a current of 250 μA was applied, and scans were conducted for a full rotation of 360°. Raw images (.tif) acquired from the scanning were reconstructed into cross-sections (.bmp) by using NRecon (SkyScan) software from the total number of 360° projection images. To check the details of the scanned specimen (externally and internally), we viewed the reconstructed image using DataViewer before we proceeded with 3D volume rendering.

2.5. Regeneration experiment

In the beginning, polychaetes were amputated at either segment four or 10 to observe the highest successful regeneration (Pires et al., 2012). We concurred that regeneration was more successful from those amputated at segment four based on preliminary study and thus proceeded with this segment for further experiments. Prior to amputation, polychaetes were removed from their tubes by a blunt end stick from the posterior opening and anaesthetised at 4% MgCl2,6H2O. Amputation was made by using a scalpel under a stereomicroscope. The posterior ends were pushed back inside the tube by forceps with the anterior part entered first. The tube was then located in a cylinder plastic container (7 cm diameter x 15 cm height) with sediments inside. The container was fully submersed in an aquarium that contained artificial aerated seawater. At this stage, they were left without food since they had no mouth. Images of successfully regenerated polychaetes at certain days after amputation until day 60 were captured.

2.6. Ultrastructure investigation

This experiment was conducted following the method provided by Imaging Center, Faculty of Pharmacy, UiTM Puncak Alam, Selangor with slight modifications. Blastema tissue appeared on day 6 p.a (D6 polychaete) was involved in this experiment. Besides blastema tissue, the investigation was also conducted on the ventral nerve cord. It was speculated that ganglion in the ventral nerve cord near the amputation site might play a crucial role in triggering NSR. Hence, two anterior-most segments from D6 polychaete, one anterior-most segment from the original body connecting with a new segment of D14 polychaete, and segments 4 to 5 from INT polychaete were amputated and fixed in 4% glutaraldehyde overnight at 4 °C. The samples were then washed three times in 0.1 M cacodylat buffer for 10 min each. Next, the samples were fixed with 1% osmium tetroxide for 2 h at 4 °C. The samples were then washed with cacodylate buffer three times for 10 min each at RT. Later, the samples were dehydrated in the series of ascending concentrations of acetone (30, 50, 75, 95% for 10 min each) and lastly 100% three times for 15 min each. The samples were then infiltrated with a series of resin: acetone mixture (resin: acetone (1 : 1); resin: acetone (3 : 1)) for 1 and 2 h respectively, followed by an overnight in resin only. Next, the resin was aspirated out, and the newly prepared resin was introduced for 2 h. The specimens were then transferred into beam capsules (later called specimen block) and filled up with resin and polymerised in an oven at 60 °C for 48 h. Next, the specimen blocks were trimmed and sectioned using an ultramicrotome (Leica EM UC7; Wetzlar, Germany). The sections were transferred onto a copper grid, stained and counterstained with 2% uranyl acetate and lead citrate for 10 min, respectively. The grids finally were examined under a transmission electron microscope (Tecnai G2 20; Oregon, USA).

2.7. Total RNA isolation

Tissues involved in RNA isolation were explained in ultrastructure experiment with n = 2 for each group. The isolation procedure was conducted according to the protocol by the NucleoSpin® RNA Plus (Macherey-Nagel, Düren, Germany). The first step involved the crushing of the tissue in liquid nitrogen using mortar and pestle. The crushed tissues were carefully inserted inside frozen-RNase-free 1.5 ml tubes and lysed with 350 μl lysis buffer. Next, the homogenised lysate was transferred to another tube in a column provided in the kit and centrifuged for 30 s at 11,000 rpm. Then, the column was discarded, and the flow-through was mixed with 100 μl binding solution by moderate vortexing. The whole lysate was transferred to another column pre-assembled with a collection tube and centrifuged for 15 s at 11,000 rpm. After that, 200 μl washing buffer #1 (provided) was added to the column and centrifuged for 15 s at 11,000 × g. The flow-through was discarded, and the column was placed in a new collection tube. Next, 600 μl of washing buffer #2 (provided) was added into the column and was centrifuged at 11,000 × g for 15 s before the step was repeated for another 250 μl washing buffer #2 that was centrifuged for 2 min at 11,000 rpm. The final step was the elution of RNA where 30 μl of RNAse-free water was added inside the previous tube and centrifuged at 11,000 × g for 1 min before the step was repeated once more. The quality of RNA was assessed on a 2100 Bioanalyzer (Agilent Inc, Santa Clara, CA, USA) and agarose gel electrophoresis. The yields were stored in -80 °C until use.

2.8. Library preparation and RNA-seq

All steps for preparing libraries and sequencing were conducted at the Malaysia Genome Institute, Malaysia by referring to TruSeq Stranded mRNA Sample Protocol (Ilumina, San Diego, CA, USA). The poly-A containing mRNA from total RNA that was previously extracted was purified by using poly-T oligo attached with magnetic beads. After that, the mRNA was fragmented into small pieces by divergent cations under elevated temperature. These fragments then were copied into the first
cDNA using reverse transcriptase and random primers. Actinomycin D was added to prevent the non-genuine DNA-dependent synthesis, and at the same time allowing RNA-dependent synthesis to improve strand specificity. The strand specificity was achieved by replacing dTTP with dUTP in the second strand marking mix and followed by second-strand cDNA synthesis using DNA polymerase 1 and RNase H. The inclusion of dUTP in the second-strand synthesis reduced the second strand during amplification. The blunt fragments tend to ligate to each other during the adapter-ligation reaction; thus, a single 'A' nucleotide was added to the 5' end of the fragments. The next step involved the enrichment of the DNA fragments by polymerase by PCR and was purified to create the final cDNA library. Finally, the cDNA libraries for each group were sequenced on an Illumina HiSeq 2000 (Illumina, San Diego, CA, USA) and data obtained were generated in FASTQ format.

2.9. Bioinformatic analysis

To avoid the generation of spurious contigs by phix-174 contamination, RNA-seq reads (FASTQ) which have forward and reverse reads for each sample, were trimmed and mapped using Bowtie 2 (Langmead and Salzberg, 2012), also known as cleaned reads. In de novo assembly, the cleaned reads were assembled using Trinity (GitHub, San Francisco, CA, USA), and genes with differential expression were identified using Empirical Analysis of Digital Gene Expression in R software (edgeR) (Bioconductor). A p-value < 0.05 was considered significantly different.
between groups. Next, the transcripts were filtered by CDHit to eliminate the redundant transcripts, and the non-redundant transcripts were blasted for protein profiles (.xml). The blasted transcripts were then loaded in Blast2GO software for mapping to retrieve GO terms, and annotation to select reliable biological processes of each protein.

2.10. Statistical analysis

Statistical analysis was only involved in the analysis of sequencing data. The differential expression of transcripts from each group was determined by pairwise comparisons using edgeR software from Bioconductor. The significant level was decided at p-value < 0.05.

3. Results

3.1. Characterisation of the anterior nervous system

For the first time, the morphology of the ANS in *D. claparedii* is fully described. Our findings from anterior regions stained by cresyl violet and labelled by anti-acetylated α-tubulin exposed the brain, branches of nerve and nerve cord. The brain is symmetrically divided into two attached hemispheres; a pair of palp nerves branching anteriorly and a pair of circumpharyngeal nerves branching posteriorly (Figure 2a). Neuron cell bodies known as perikarya (sing., perikaryon) were accumulated around the brain with nerve fibre mass in the middle, known as neuropil (Figure 2b). The double-stranded nerve cord was observed along the ventral side with a pair of segmental ganglia connected by commissure. In addition, ganglion from each segment was connected by a connective (Figure 2c). All these structures form the CNS of *D. claparedii*.

Our finding from anti-acetylated α-tubulin labelling confirmed the nerve structures as mentioned above with additional nuchal organs, nuchal nerves and lateral nerves of PNS (Figure 2d). Interestingly, the nerve observed under confocal laser scanning microscope also revealed an additional median nerve in the nerve cord, suggesting the possibility of *Diopatra claparedii* has a triple-stranded nerve cord (Figure 2e). Furthermore, we conducted micro-computed tomography scanning (micro-CT) to confirm the location of these nerve structures in *D. claparedii*. *D. claparedii* brain is located at the dorsal region of the head known as the prostomium, just above the pharynx. Meanwhile, the nerve cord is located ventrally started at around 7th segment (Figures 2f, 2g).

3.2. Cellular evidence of early anterior NSR in *D. claparedii*

Prior to the cellular investigation of early NSR, we observed the full anterior regeneration after the polychaete was amputated at the 4th segment (Figure 2). On day 1 post-amputation (p.a.), the amputated site was closed without infection (Figure 3a). We also noticed the appearance of tissue mass at the middle body plane of the amputation site called blastema at day 6 p.a. (Figure 3b). Later on day 14 p.a., we observed new anterior (prostomium and peristomium) appeared, somewhat translucent. The new anterior arose in the middle of the amputation site where the gut is located (Figure 3c). On day 30 p.a., segmentation is seen

![Figure 3](image-url)

**Figure 3.** Anterior regeneration in *D. claparedii*. (a) Ventral view. A recovering polychaete at day 1 p.a. Perforated line is where the polychaete was amputated. (b) Axial view. Condition at day 6 p.a. with the presence of blastema tissue (blas) at the mid body plane. (c) Ventral view. Anterior regeneration at day 14 p.a. New head formation with three antennae. (d) Lateral view. Polychaete regenerating at day 30 p.a. Segmentation is obvious as marked by the double-headed arrow. Five new segments arose from four losing segments. (e) Ventral view. Almost complete regeneration at day 60 p.a. Branchiae (bran) was smaller compared to the old body. The mouth has developed and the feeding behaviour has become normal. New segments arose more than the ones amputated. Scale bars = 1 mm.
replacing the lost segments. However, our observation found that new segments were not replacing the exact number of lost segments (Figure 3d). The full anterior regeneration was observed at around day 60 p.a. by comparing the new anterior width with the original body. The widths of the new anterior and the original body were almost the same. However, in the new anterior, branchiae were smaller, and the overall colour of the new anterior was lighter than the original body.

The ultrastructure analyses on NSR were conducted on tissues of three groups; ganglion in the intact polychaete (INT), ganglion and blastema in polychaete at day 6 p.a (D6) and ganglion in polychaete of day 14 p.a. (D14). Electron micrograph (Figure 4a) clearly shows the synaptic transmissions between axon terminals and dendrites in the INT ganglion, specifically the neuropil, indicating the occurrence of a neurophysiology process. Furthermore, we also confirmed the presence of perikarya or ganglionic cells from the intact ganglion (Figure 4a). Our TEM findings also revealed distinctive differences between intact and regenerating polychaetes. Compared to INT polychaete, the presence of lysosomes and lipofuscins in both D6 and D14 polychaetes were very apparent, indicating that the cell digestions process was underway (Figure 4b).

3.3. Molecular evidence of early NSR in D. claparedii

We investigated the NSR in our polychaete species by conducting RNA-seq on Illumina sequencer. The tissues from the three regenerating stages, i.e. INT, D6 and D14 (n = 2 each group stage) were extracted for their mRNAs prior to their conversion to six libraries. Phix-174 was used as quality and calibration control for Illumina sequencing. Each sequence generated from Illumina sequencing platform was screened against phix-174 to remove the contaminants. RNA-seq analysis has identified 147,445 differential expressed genes (transcripts) with p < 0.05. The

Figure 4. The occurrence of NSR analysed by TEM. (a) Ultrastructures in INT ganglion. Left image: Neuropil showing synaptic transmission between axon terminals (yellow areas) and dendrites (green areas). The transmissions were seen at the synaptic clefts (red arrows). Synaptic vesicles (s.v) are located inside the axon terminals. Right image: Accumulation of perikarya (also from semithin section seen in insert). The nucleus (nuc) is always seen as a prominent structure in a neuron. Perikaryon also performs synaptic connectivity with axon terminal since somatic vesicles (s.o.v) containing neurotransmitter also present there. (b) The appearance of lysosomes in D6 and D14. Left image: Accumulation of lysosomes (white arrows) in blastema cell of D6 polychaete. Middle image: Final stage of lysosomal activity seen in ganglion of D6 polychaete. Right image: Lysosomes (white arrows) in the ganglion of D14 polychaete. (c) Differences in the shape of mitochondria (red arrows) between D6 and D14 polychaetes. Note as well both polychaetes have multiple nucleoli (ni). Left image: Blastema cells (in D6) are recognised by having numerous mitochondria indicating high synthetic activity in the cells. The shape of mitochondria were rounded. Right image: Mitochondria in D14 were elongated in shape and numerous.
The average contig length was 890.58 bp. Figure 5a shows differentially expressed transcripts (red dots) when compared between groups with \( p < 0.05 \). Moreover, Figure 5b shows differentially expressed transcripts between groups with \( p < 1 \times 10^{-10} \), indicating up- and down-regulation. The transcripts were filtered using CDHit, and 116,203 non-redundant transcripts were obtained. These transcripts were then predicted for protein sequences by Transdecoder, and a total of 37,248 coding sequences were recorded.

3.4. Differential expression of proteins related to NSR

The analysis conducted towards transcriptome in the regenerating anterior of *D. claparedii* has identified multiple proteins. Gene Ontology (GO) mapping was conducted to retrieve candidate GO terms associated with the hits obtained after a BLAST search. Then, the proteins were mapped by using Blast2GO (BioBam, Valencia, Spain) to obtain the functional labels associated with the processes that occur in the extracted tissues. Specifically for NSR, GO terms (related to biological function) were listed as i) neurogenesis, ii) nervous system development, iii) central nervous system development, iv) neuron migration, v) regulation of neurogenesis, vi) axon regeneration, vii) positive regulation of axon regeneration, viii) axon extension, ix) regulation of axon extension, x) neuron projection regeneration, xi) axonogenesis, xii) neuron development, xiii) axon development, xiv) generation of neurons, xv) neuron differentiation, xvi) peripheral nervous system development. The sequences obtained have been submitted to NCBI (Table 1).

Table 2 summarises the proteins expressed during nervous system regeneration based on the GO terms listed, which are significantly different \( (p < 0.05) \) compared between groups and exist at \( -2 > \text{logFC} > 2 \). The highest protein expressed and upregulated during NSR in *D. claparedii* was noelin-like isoform X2, which categorised under nervous system development (GO:0007399). The protein was differentially expressed in both D6 and D14 polychaetes compared to INT. In contrast, smoothened homolog was highly downregulated in both D6 and D14 compared to INT based on its logFC. This protein was categorised under GO:0031102, i.e. neuron projection regeneration. Also, several proteins...
Table 1. Sequencing data accession numbers (https://dataview.ncbi.nlm.nih.gov/object/PRJNA663357?reviewer=i8oa2u3reioo342o8v9gnf5ahc).

| No. | Accession No. | Title | BioProject | BioSample | Library ID |
|-----|---------------|-------|------------|-----------|------------|
| 1   | SRR12672111   | Intact segment 4 to 5 of anterior body: replicate 1 | PRJNA663357 | SAMN16131181 | INT-1 |
| 2   | SRR12672110   | Intact segment 4 to 5 of anterior body: replicate 2 | PRJNA663357 | SAMN16131182 | INT-2 |
| 3   | SRR12672109   | Day 6 post amputation replicate 1 | PRJNA663357 | SAMN16131183 | D6-1 |
| 4   | SRR12672108   | Day 6 post amputation replicate 2 | PRJNA663357 | SAMN16131184 | D6-2 |
| 5   | SRR12672107   | Day 14 post amputation replicate 1 | PRJNA663357 | SAMN16131185 | D14-1 |
| 6   | SRR12672106   | Day 14 post amputation replicate 2 | PRJNA663357 | SAMN16131186 | D14-2 |

Table 2. Selected Genes Ontology categories represented along the anterior NSR. All differences significant compared to INT, except * significant when compared with D6.

| GO Term                          | Description                        | Protein Name | Group (n = 2) (P value) | Up-regulation | Down-regulation |
|----------------------------------|------------------------------------|--------------|-------------------------|---------------|-----------------|
| GO:0007399 nervous system development | noelin-like isoform X2             | D6 (2.72E-23) D14 (2.12E-23) | | |
| | polycomb complex protein BMI-1-B-like isoform X2 | D6 (6.23E-14) D14 (1.54E-07) | | |
| | polycomb complex protein BMI-1 | D6 (9.21E-13) D14 (1.29E-08) | | |
| | trafficking protein particle complex subunit 6B | D6 (1.34E-09) D14 (6.69E-08) | | |
| | noelin-like isoform X3 | D14 (8.84E-07) *D14 (0.0002779) | | |
| | proteoglycan 4 | D6 (1.83E-08) *D14 (1.72E-06) | | |
| | protein unc-119 homolog B-like | D6 (9.48E-06) *D14 (1.45E-05) | | |
| | protocadherin alpha-11-like | D14 (0.0196149) | | |
| | zinc finger protein 335 isoform X1 | D6 (0.0010263) | | |
| | glycoprotein 3-alpha-L-fucosyltransferase A | D14 (0.0001255) | | |
| GO: 0031102 neuron projection regeneration | smoothed homolog | D14 (1.66E-12) D6 (9.62E-09) | | |
| GO:0007417 central nervous system development | low-density lipoprotein receptor-related protein 4-like | D14 (1.83E-11) | | |
| | hyaluronan and proteoglycan link protein 3-like | D6 (9.30E-05) D14 (0.0220912) | | |
| | versican core protein-like | D6 (2.05E-06) D14 (8.19E-05) | | |
| | hyaluronan and proteoglycan link protein 1-like | D14 (0.0124250) | | |
| | achaete-scute homolog 1 | D14 (0.0004928) | | |
| | versican core protein | D14 (0.0001256) | | |
| | transcription factor SOX-8 | D14 (0.0001256) | | |
| GO:0022008 neurogenesis | neuron navigator 2 isofrom X9 | D14 (0.0005557) *D14 (6.40E-09) D14 (0.0004403) | | |
| | BTB/POZ domain-containing protein 6-like | D14 (1.12E-07) | | |
| | transcription factor SOX-8 | *D14 (7.45E-05) D14 (0.0001256) | | |
| GO: 0007409 axonogenesis | cartilage matrix protein | D14 (2.75E-08) *D14 (1.19E-05) | | |
| | slit homolog 1 protein | D6 (4.00E-07) D14 (3.45E-07) | | |
| | immunoglobulin A1 protease | D14 (8.70E-05) D6 (0.0085116) | | |
| | cartilage matrix protein | D14 (0.0043983) *D14 (0.000296) | | |
| GO: 0048699 generation of neurons | protein IMPACT isoform X1 | D14 (4.28E-06) | | |
| GO:0050767 regulation of neurogenesis | EGF-like domain-containing protein 1 | D14 (3.47E-07) *D14 (3.23E-05) | | |
| | microfibril-associated glycoprotein 4-like isoform X1 | D6 (5.43E-07) | | |
| GO: 0030182 neuron differentiation | receptor-type tyrosine-protein phosphatase delta-like | D6 (0.001185) | | |
| | protein Wnt-16-like | D6 (2.23E-07) | | |
| | protein Wnt-5b-like | D14 (7.45E-05) *D14 (4.53E-05) | | |
| GO: 0048666 neuron development | achaete-scute homolog 1 | D14 (0.0039118) D14 (0.0124250) | | |
allow us to distinguish other nerve branches as seen in the conventional histology, including the combination of immunohistochemistry and confocal laser scanning microscopy (IHC + CLSM). The inability of micro-CT to distinguish nerve branches were also reported in several studies (Faulwetter et al., 2013; Parapar et al., 2017). Images of soft tissue such as nerves are usually not well recorded due to low intrinsic X-ray contrast on non-mineralised tissue (Thompson et al., 2020).

Through TEM, perikaryon was observed to contain the prominent nucleus, numerous mitochondria, and synaptic vesicles. The presence of neuropil indicated a complex connection of synapses between axon terminals and dendrites, enabling synaptic transmissions to occur. The vesicles of neurotransmitters or neuropeptide were also observed, suggesting the effect of vesicular volume on exocytosis (Sombers et al., 2004). The observation is similar to the previous study on annellar regeneration from a cellular perspective (Bely, 2014). However, previous study was limited up to blastema formation, while in the present study, we extended this to the early segmentation stage, which was seen in D14 polychaete. For a regeneration process to occur, blastema should start to redifferentiate into specific structures (Bely, 2014). Our findings from TEM magnified the ultrastructures that appeared in blastema tissue. The apparent appearance of lysosomes was seen by electron-dense materials. Lysosome consists of degradative enzymes that break down unwanted biomolecules during cellular metabolism, including regeneration. Furthermore, lysosomes also store Ca^{2+}, which can be released into the cytoplasm for intracellular functions (Hesketh et al., 2018). Numerous mitochondria were also apparent in both D6 and D14 polychaetes; however, their shapes changed along the time. Mitochondria in D6 polychaete was round, while the ones in D14 were elongated. A round-shaped mitochondrion can produce more reactive oxygen species (Ahmad et al., 2013), which involves in apoptosis during the wound healing process (Bryan et al., 2012; Cano Sanchez et al., 2018). The presence of elongated mitochondria indicate protection from degradation. The mitochondria have more cristae, meaning increased ATP synthase activity, thus maintaining the cells’ viability to continue regenerating. This condition is known as a cellular stress response (Fulda et al., 2010). Another indication that cells were actively regenerating was the appearance of multiple nucleoli in the nucleus (Pedersen, 1972; Maki et al., 2007). Nucleolus composed of filamentous and granular structures and production site of ribosomal RNA. The RNA then will produce ribosomal protein to become ribosomal subunit (Eeken, 2001), which is crucial in producing specific proteins for regeneration.

The transcriptomes of the NSR in amputated D. claparedii in two regenerating stages, as well as intact polychaetes, were sequenced and compared to each other. Through this experiment, we have identified proteins involved in NSR and other processes related to the production of new nerve tissues. Our finding has identified nine categories from GO terms that satisfied the significant level (p < 0.05) and fold change (≥ 2 & ≤ -2). Under the nervous system development (GO: 0007399), five proteins were upregulated. The highest differentially expressed and highest fold change belonged to the noelin-like isoform X2, a putative protein for noelin (OLF1M). We also found that noelin-like isoform X3 is among the highest upregulated protein that differentially expressed. Noelin or olfactomedin (Anholt, 2014) belongs to olfactomedin domain-containing proteins and is involved in neuronal differentiation during development, neurogenesis, and generation of neural crest during embryogenesis and larval development (Moreno and Bronner-Fraser, 2005; Rice et al., 2012). Olfactomedin 1 was also reported to suppress the cleavage of amyloid precursor protein, inhibit the subsequent production of amyloid-beta, the main contributor to Alzheimer’s disease (Takahama et al., 2004). Furthermore, slt homolog 1 protein was also expressed in the regenerating polychaetes. This protein is reportedly involved in axon guidance or navigation during cellular migration (Wong et al., 2002). This protein has not been documented anywhere except in an article by Ribeiro et al. (2019) who studied the anterior regeneration in a syllid worm. Protein unc-119 homolog B-like is similar to unc-119 that functioning in neural architecture patterning. This
protein is well conserved in metazoa, and the mutation to this protein will lead to nervous system disorientation and paralysis (Manning et al., 2004). However, this protein was downregulated in D14 polychaete that might show neural architecture patterning has been completed. The last relevant protein in NSR is neuron navigator 2 isoform X9 which might show neural architecture patterning has been completed. The last relevant protein in NSR is neuron navigator 2 isoform X9 which might be involved in cell migration and axonal outgrowth (Maes et al., 2002; Muley et al., 2008) and it was significantly expressed in D14 which probably true since the nervous system was in the process of development.

5. Conclusions

In summary, we have presented the characteristics of the anterior nervous system in D. clapareedi morphologically, i.e. the region for anterior NSR. We have highlighted the brain and ventral nerve cord as the main structures in CNS. We further investigated the ultrastructures, particularly the ganglion of the nerve cord from the regenerating intact tissues and concluded significant differences between the regenerating stages. Our molecular data also revealed new putative proteins responsible in nervous system regeneration requiring further extensive investigation as to further experimentation will provide comprehensive information on questions such as real-time protein activities during nervous system regeneration.

Declarations

Author contribution statement

Mohd Ulil Ilmie Ahmad Nazri, Wan Iryani Wan Ismail: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Mohd Hafizi Mahmud, Basri Saidi, Mohd Noor Mat Isa, Zolkapli Ehsak: Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Othman Ross: Conceived and designed the experiments; Wrote the paper.

Izwandy Idris: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Funding statement

This work was supported by the Ministry of Higher Education (MOHE), FRGS/1/2016/WAB09/UMT/02/2.

Data availability statement

Data associated with this study has been deposited at NIH under the url https://dataview.ncbi.nlm.nih.gov/object/PRJNA663357?reviewer=i8oa2u3reioo342o8v9gnf5ahc.

Declaration of interests statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

Acknowledgements

We would like to acknowledge Professor Pat Hutchings from Australian Museum Research Institute, Sydney, Australia for proof-reading the manuscript. Special thanks to Senec Bol from Mah Meri tribe in Morib for helping us with the specimen collection.

References

Ahmad, T., et al., 2013. Computational classification of mitochondrial shapes reflects stress and redox state. Cell Death Dis. 4, e641.

Aliardi, L., 2009. Morphological and Cellular Aspects of Tail and Limb Regeneration in Lizards: A Model System with Implications for Tissue Regeneration in Mammals. Springer Science & Business Media.

Anderton, D.T., 2013. Embryology and phylogeny in annelids and arthropods: international series of monographs in pure and applied biology zoology. In: International Series of Monographs on Pure and Applied Biology: Division, Zoology, Vol. 50. Elsevier.

Anholt, R., 2014. Olfactomedin proteins: central players in development and disease. Front. Cell. Dev. Biol. 2, 6.

Bely, A.E., 2014. Early events in anemid regeneration: a cellular perspective. Integr. Comp. Biol. 54, 688–699.

Brinkmann, N., Wanninger, A., 2009. Neurogenesis suggests independent evolution of opercula in serpulid polychaetes. BMC Evol. Biol. 9, 270.

Bryan, N., et al., 2012. Reactive oxygen species (ROS)—a family of fate deciding molecules pivotal in constitutive inflammation and wound healing. Eur. Cell. Mater. 24, e65.

Cano Sanchez, M., Lancel, S., Boulanger, E., Neviere, R., 2018. Targeting oxidative stress and mitochondrial dysfunction in the treatment of impaired wound healing: a systematic review. Antioxidants 7, 98.

Cayre, M., Malaterrre, J., Scotto-Lussanne, S., Strambi, C., Strambi, A., 2002. The common properties of neurogenesis in the adult brain: from invertebrates to vertebrates. Comp. Biochem. Physiol. Part B Biochem. Mol. Biol. 132 (1), 1–15.

Eeken, J.C., 2001. J. Nucleolus. Elsevier.

Eckert, M., et al., 2011. Neuroregeneration in neurodegenerative disorders. BMC Neuro. 11, 75.

Faulwetter, S., Vasileiadou, A., Kouratoras, M., Dailianis, T., Arvanitidis, C., 2013. Micro-computed tomography: introducing new dimensions to taxonomy. Zookeys 263, 1–45.

Fulda, S., Gorman, A.M., Hori, O., Samali, A., 2010. Cellular stress responses: cell survival and cell death. Int. J. Cell Biol. 2010, 1–23.

Gemberling, M., Bailey, T.J., Hyde, D.R., Pons, K.D., 2013. The zebrafish as a model for complex tissue regeneration. Trends Genet. 29, 611–620.

Helm, C., et al., 2018. Convergent evolution of the ladder-like ventral nerve cord in Annelida. Front. Zool. 15, 36.

Hesketh, G.G., Wartosch, L., Davis, L.J., Bright, N.A., Luzio, J.P., 2018. The lysosome and intracellular signalling. In: Lamas, C., Prior, I. (Eds.), Endocytosis and Signalling Progress in Molecular and Subcellular Biology, ume 57, Springer, pp. 151–180.

Idris, I., Anshad, A., 2013. Checklist of polychaetous annelids in Malaysia with redescriptions of two commercially exploited species. Asian J. Anim. Vet. Adv. 8, 1–43.

Mackie, A.S., 1994. Collecting and preserving polychaetes. Polychaete Res. 16, 7–9.

Maes, T., Barcelo, A., Buesa, C., 2002. Neuron navigator: a human gene family with homology to unc-53, a cell guidance gene from Caenohabditis elegans. Genomics 80 (1), 21–30.

Maki, N., et al., 2007. Rapid accumulation of nucleostemin in nucleolus during newt nervous system regeneration. Dev. Dyn. 236, 941–955.

Manning, A.G., Crawford, B.D., Waskiewicz, A.J., Pilgrim, D.B., 2004. UNC-19 homolog required for normal development of the zebrafish nervous system. Genetics 40 (4), 223–230.

Moreno, T.A., Bronner-Frazer, M., 2005. Noelin modulates the timing of neuronal differentiation during development. Dev. Biol. 288, 434–447.

Muley, P.D., McNicol, E.M., Marzinke, M.A., Knobel, K.M., Barr, M.M., Clagett-Dame, M., 2008. The aVRa-responsive gene neuron navigator 2 functions in neurite outgrowth and axonal elongation. Develop. Neurobiol. 68 (13), 1441–1453.

Müller, M.C., Berenzen, A., Westbridge, W., 2003. Experiments on anterior regeneration in Eurythoe complanata (‘Polychaeta’, Amphipodiinae): reconfiguration of the nervous system and its function for regeneration. Zoomorphology 122, 95–103.

Nazri, M.U.I.A., Idris, I., Ross, O., Ismail, W.I.W., 2019. Neurological disorder brain model: a lesson from marine worms (annelida: polychaeta). Malays. J. Med. Sci. 26, 5–18.

Orthage, L., 1995. On the innervation and homologies of the anterior end appendages of the Eunicea (Polychaeta), with a tentative outline of the fundamental constitution of the cephalic nervous system of the polychaeta. Acta Zool. 76, 229–248.

Ortega, A., Oliveras-Baneuelt, T.N., 2020. Neurons and glia cells in marine invertebrates: an update. Front. Neurosci. 14, 121.

Parapar, J., Candás, M., Carda-Beiras, X., Moreira, J., 2017. Exploring annelid anatomy using micro-computed tomography: a taxonomic approach. Zool. Anz. 270, 19–42.

Pedersen, K.J., 1972. Studies on regeneration blastemans of the planarian Dugesia tigrina with special reference to differentiation of the muscle-connective tissue filament system. Wilhelm Roux. Arch. Entwickl. Mech. Org. 169, 134–158.

Purves, K., 2005. Sense organs in polychaetes (Annelida). In: Purschke (Ed.), Morphology, Molecules, Evolution and Phylogeny in Polychaeta and Related Taxa. Springer, pp. 53–78.

Fires, A., Freitas, R., Quintino, V., Rodrigues, A.M., 2012. Can dismutase neopoliudina (annelida: enoploidae) regenerate body damage caused by bait digging or predation? Estuar. Coast Shelf Sci. 110, 36–42.

Pyza, E.M., 2013. Plasticity in invertebrate sensory systems. Front. Physiol. 4, 226.

Reuter, M., et al., 1996. Development of the nervous system in Dugesia tigrina during regeneration after finotom and decapitation. Invertebr. Reprod. Dev. 29, 199–211.

Ribeiro, R.P., Fonseca, A., Biedron, C., Agudo, M.T., 2019. Comparative transcriptomics in Syllidae (Annelida) indicates that posterior regeneration and regular growth are comparable, while anterior regeneration is a distinct process. BMC Genom. 20 (1), 855.
Rice, H.C., et al., 2012. Pancortins interact with amyloid precursor protein and modulate cortical cell migration. Development 139, 3986–3996.
Rouse, G.W., Fauchald, K., 1997. Cladistics and polychaetes. Zool. Sci. 26, 139–204.
Rouse, G., Pleijel, F., 2001. Polychaetes. Oxford University Press.
Sombers, L.A., et al., 2004. The effects of vesicular volume on secretion through the fusion pore in exocytotic release from PC12 cells. J. Neurosci. 24, 303–309.
Takahama, S., Nakaya, N., Tomarev, S.I., 2004. Olfactomedin 1 may suppress APP cleavage through its interaction with BACE1. Invest. Ophthalmo. Vis. Sci. 55, 2959-2959.
Thorp, J.H., Rogers, D.C., 2014. Thorp and Covich’s Freshwater Invertebrates: Ecology and General Biology, ume 1. Elsevier.
Thompson, N., et al., 2020. MicroCT optimisation for imaging fascicular anatomy in peripheral nerves. J. Neurosci. Methods 338, 108652.
Weidhase, M., Helm, C., Bleidorn, C., 2015. Morphological investigations of posttraumatic regeneration in Timarete cf. punctata (Annelida: cirratulidae). Zool. Lett. 1, 1–16.
Wong, K., Park, H.T., Wu, J.Y., Rao, Y., 2002. Slit proteins: molecular guidance cues for cells ranging from neurons to leukocytes. Curr. Opin. Genet. Dev. 12 (5), 583–591.
Yoshida-Noro, C., Myohara, M., Kobari, F., Tochinai, S., 2000. Nervous system dynamics during fragmentation and regeneration in Enchytraeus japonensis (Oligochaeta, Annelida). Dev. Gene. Evol. 210, 311–319.