



Report on Current and Future Biological and Ecosystem Science Research Activities

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PURPOSE

To provide the Scientific Review Board with a description of progress towards research activities described in the IPHC's five-year Program of Integrated Research and Monitoring (2022-2026).

BACKGROUND

The primary biological and ecological research activities at the IPHC that follow Commission objectives are identified and described in the IPHC Five-Year Program of Integrated Research and Monitoring (2022-2026). These activities are integrated with stock assessment (SA) and the management strategy evaluation (MSE) processes (Appendix I) and are summarized in five main areas, as follows:

- 1) Migration and Population Dynamics. Studies are aimed at improving current knowledge of Pacific halibut migration and population dynamics throughout all life stages in order to achieve a complete understanding of stock structure and distribution across the entire distribution range of Pacific halibut in the North Pacific Ocean and the biotic and abiotic factors that influence it.
- 2) Reproduction. Studies are aimed at providing information on the sex ratio of the commercial catch and to improve current estimates of maturity and fecundity.
- 3) Growth. Studies are aimed at describing the role of factors responsible for the observed changes in size-at-age and at evaluating growth and physiological condition in Pacific halibut.
- 4) Mortality and Survival Assessment. Studies are aimed at providing updated estimates of discard mortality rates in the guided recreational fisheries and at evaluating methods for reducing mortality of Pacific halibut.
- 5) Fishing Technology. Studies are aimed at developing methods that involve modifications of fishing gear with the purpose of reducing Pacific halibut mortality due to depredation and bycatch.

A ranked list of biological uncertainties and parameters for SA (Appendix II) and the MSE process (Appendix III) and their links to research activities and outcomes derived from the five-year research plan are provided.

SRB REQUESTS AND RECOMMENDATIONS

The SRB issued the following recommendation in their report of SRB028 (IPHC-2025-SRB028-R) in relation to presentation IPHC-2026-SRB028-06:

*SRB028–Req.01 ([para. 16](#)) The SRB **REQUESTED** an analysis that separates year and biological region effects on maturity ogives. This would answer whether variation in age at maturity is driven by variation in the spatial distribution of the stock or by interannual variation in environmental conditions.*

The IPHC Secretariat has conducted analyses of histology-based maturity data separating year and biological region using best-fit logistic GAM. These data were provided in IPHC-2025-SRB026-06. Potential drivers of variation in age at maturity are being investigated.

SRB028–Req.02 ([para. 17](#)) The SRB **REQUESTED** that the Secretariat investigate:

- a) *environmental predictors of interannual variation in maturity ogives; and*
- b) *the potential to include time-varying maturity ogives in the stock assessment.*

The IPhC Secretariat is investigating potential predictors of interannual variation in maturity ogives during the period of time (2002-2024) during which visual maturity was calibrated by applying the calibration using histological and visual data (2022-2024).

SRB028–Req.03 ([para. 18](#)) The SRB **REQUESTED** that, as more epigenetic aging data become available, a more systematic approach to cross-validation be developed in which performance is evaluated across space and time. For example, applying a model trained on samples from all but one area to the withheld area and applying a model trained on samples from all but one year to the withheld year. Ideally, a full factorial approach could be used to determine whether any training-test combinations perform inadequately.

The IPhC Secretariat is investigating approaches to cross-validate epigenetic data across sampled areas (space) and (years) (time).

SRB028–Rec.02 ([para. 35](#)) **NOTING** that the Secretariat has developed multiple aging methods (epigenetic, AI-based, traditional break and burn otolith aging), the SRB **RECOMMENDED** that a table be developed – with a column for each method - describing:

- a) *the different anticipated use cases;*
- b) *the relative costs (split into development and production) and benefits of these approaches;*
- c) *timelines for development; and*
- d) *decision points that will determine future allocation of effort to this method.*

The IPhC Secretariat has incorporated information related to epigenetic aging into a table that includes multiple aging methods for comparison.

UPDATE ON PROGRESS ON THE MAIN RESEARCH ACTIVITIES

1. Migration and Population Dynamics.

The IPhC Secretariat is currently focusing on studies that incorporate genomics approaches in order to produce useful information on population structure, distribution and connectivity of Pacific halibut. The relevance of research outcomes from these activities for stock assessment (SA) resides (1) in the introduction of possible changes in the structure of future stock assessments, as separate assessments may be constructed if functionally isolated components of the population are found (e.g. IPhC Regulatory Area 4B), and (2) in the improvement of productivity estimates, as this information may be used to define management targets for minimum spawning biomass by Biological Region. These research outcomes provide the second and third top ranked biological inputs into SA (Appendix II). Furthermore, the relevance of these research outcomes for the MSE process is in biological

parameterization and validation of movement estimates, on one hand, and of recruitment distribution, on the other hand (Appendix III).

- 1.1. Population genomics. The primary objective of these studies is to investigate the genetic structure of the Pacific halibut population and to conduct genetic analyses to inform on Pacific halibut population dynamics and distribution within the Convention Area

- 1.1.1. Population genetic structure. Details on sample collection, sequencing, bioinformatic processing and proposed analyses utilizing low-coverage whole genome sequencing (lcWGR) to investigate Pacific halibut population structure were provided in documents IPHC-2021-SRB018-08, IPHC-2022-SRB021-09, IPHC-2023-SRB022-09, IPHC-2024-SRB024-09 and IPHC-2025-SRB026-06. Results from these studies are currently being prepared for publication in a leading peer-reviewed journal.

- 1.1.2. Development of a novel method for estimating genetic differentiation from genotype likelihoods. As part of IPHC's research on population genetic structure, the IPHC Secretariat has developed a bioinformatic method (named *fst-gl*) designed to estimate F_{ST} using low-coverage whole genome resequencing (lcWGR) data from multiple populations. F_{ST} (Weir and Cockerham, 1984) is a widely used measure of population differentiation that is applied in the identification of SNPs or localized regions of the genome that show high levels of differentiation for the purposes of SNP panel development, identification of genomic signals of natural selection or local adaptation, identification of sex-associated genomic regions (Pacific halibut: Jasonowicz et al. 2022), etc.

Current available bioinformatic methods for estimating F_{ST} require hard-called genotypic data generated at high-coverages ($> 10x$) using methods such as restricted site associated DNA (RAD)-seq or whole genome sequencing (WGS) to obtain accurate estimates of F_{ST} . When sequencing at lower coverages ($< 5x$), uncertainty arises at the level of individual genotypes; therefore, accounting for this uncertainty is necessary when analyzing lcWGR data. The estimation of F_{ST} from lcWGR data or genotype likelihoods is complicated by the very limited number of available bioinformatic methods (e.g. *angsd*) that only estimate pairwise F_{ST} between two populations from low-coverage sequencing data (Korneliussen et al. 2014; Rasmussen et al. 2022) at a high computational cost. The developed bioinformatic method (*fst-gl*) enables the estimation of F_{ST} among any number of populations and implements resampling routines to provide bootstrapped significance values for estimates of F_{ST} , as previously detailed in document IPHC-2026-SRB028-06. A manuscript describing the development and validation of this new method has been submitted for publication to the journal *Bioinformatics* (Jasonowicz and Planas, in review).

- 1.2. Genomics-based method for estimating age of Pacific halibut. The primary objective of this project is to develop a genetic method for aging Pacific halibut using fin tissue, a sample that can be easily collected from either live or dead individuals. This method is based on the identification of DNA methylation patterns in fin tissue that are associated with age through the development of an age estimation model (i.e., an epigenetic clock)

for Pacific halibut. The first epigenetic clock was developed for humans in 2013 (Horvath, 2013), and it predicted age with great accuracy ($r = 0.96$) and with a mean aging error (MAE) of 3.6 years. Subsequently, epigenetic clocks have been developed for several fish species that demonstrated improved accuracy (r between 0.84 and 0.99) and lower average MAE (0.87 years, or 3.5% of the total lifespan of the species examined) (reviewed in Piferrer and Anastasiadi, 2023).

Patterns of DNA methylation (i.e. a natural process of regulation of gene expression that consists in the covalent modification of the nucleobase cytosine) in Pacific halibut will be investigated by performing genome-wide DNA methylation at single base-pair resolution using reduced representation bisulfite sequencing (RRBS) by leveraging the high-quality genome assembly available for Pacific halibut (Jasonowicz et al. 2022). This is an efficient and cost-efficient method to identify methylation patterns (i.e., CpG sites) in DNA because it targets bisulfite sequencing to a well-defined set of genomic regions with high CpG density that can be sequenced at high read depth. Age-associated DNA methylation patterns will be modelled to generate an epigenetic age predictor (i.e., epigenetic clock) for Pacific halibut constructed using elastic net penalized regression models that select a group of CpG sites that have a monotonically increasing relationship with age in the selected training data set. By implementing these linear models that select and weight age-correlated CpG sites, chronological age of Pacific halibut will be estimated based on the percentage methylation at these key CpG sites in fin tissue samples.

1.2.1. Methods.

1.2.1.1. Genetic samples used in the development of the epigenetic clock. For developing an epigenetic clock, we selected genetic samples (fin clips) from 249 individual Pacific halibut collected during IPHC's Fishery Independent Setline Survey (FISS) from 2021 to 2024 throughout IPHC Convention Waters (Figure 1) and covering most of IPHC's Regulatory Areas (Table 1). These genetic samples correspond to fish with known ages (read twice by the traditional break and bake aging method) between 6 to 30 years and include 6-10 individual samples (aiming at equal number of males and females) per year of age (Figure 2).

Table 1. Number of FISS samples by year and IPHC Regulatory Area with RRBS sequence data generated for the development of the Pacific halibut epigenetic clock.

Year	2A	2B	2C	3A	3B	4A	4B	4D	Total
2021	1	12	12	3	0	3	1	0	32
2022	2	22	17	20	2	2	3	1	69
2023	16	32	25	14	10	0	0	0	97
2024	0	15	26	7	1	0	0	2	51
Total	19	81	80	44	13	5	4	3	249

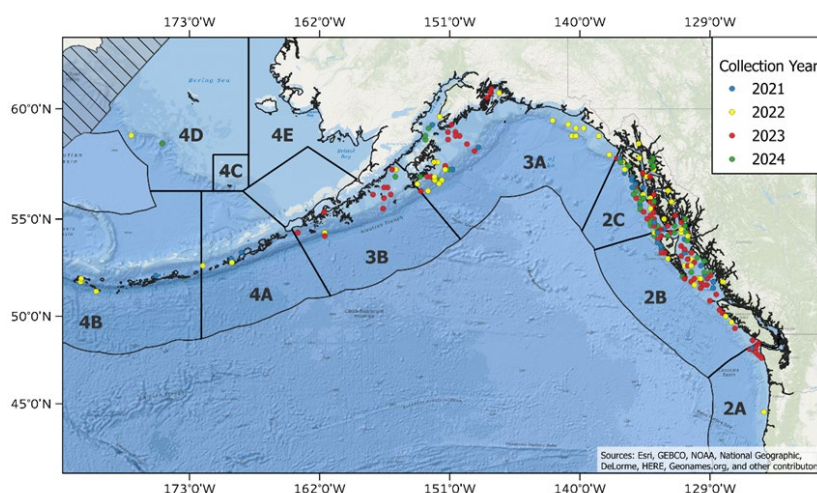


Figure 1. Map showing the collection locations of the samples with RRBS sequence data generated for the development of the Pacific halibut epigenetic clock.

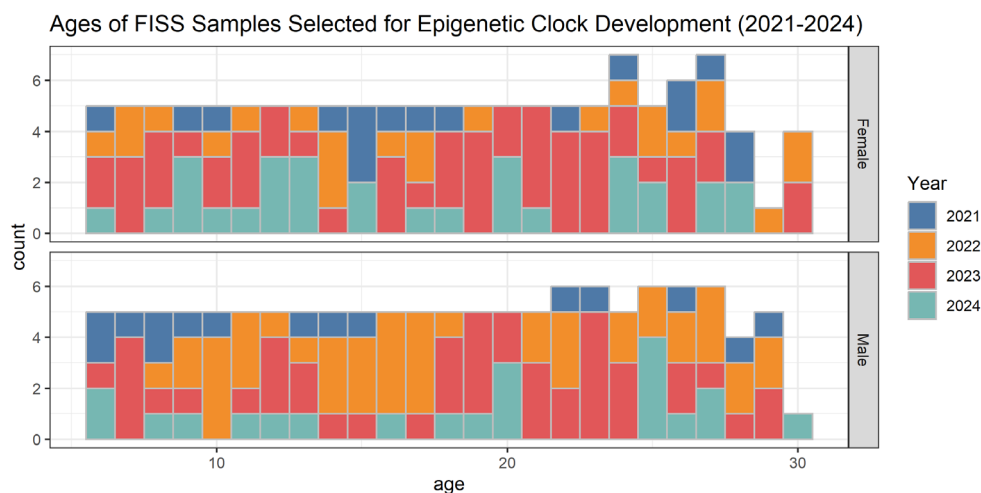


Figure 2. Histograms showing the number of female (top) and male (bottom) genetic samples with RRBS sequence data generated for the development of the Pacific halibut epigenetic clock.

1.2.1.2. Genetic samples used in testing out-of-sample interannual predictive performance.

For testing out-of-sample interannual predictive performance of the epigenetic clock, we selected genetic samples (fin clips) from 46 individual Pacific halibut collected in commercial landings from 2017 to 2020 covering most of IPHC's Regulatory Areas (Table 2). These genetic samples correspond to fish with known ages (read twice by the traditional break and bake aging method) between 6 to 27 years (Figure 3).

Table 2. Number of market samples by year and IPHC Regulatory Area submitted for RRBS sequencing on the Aviti sequencing platform for out-of-sample validation of the Pacific halibut epigenetic clock.

Year	2C	3A	3B	4A	4B	4C	4D	Total
2017	0	0	0	4	4	0	4	12
2018	1	3	2	0	5	0	1	12
2019	0	3	2	3	2	1	0	11
2020	3	4	4	0	0	0	0	11
Total	4	10	8	7	11	1	5	46

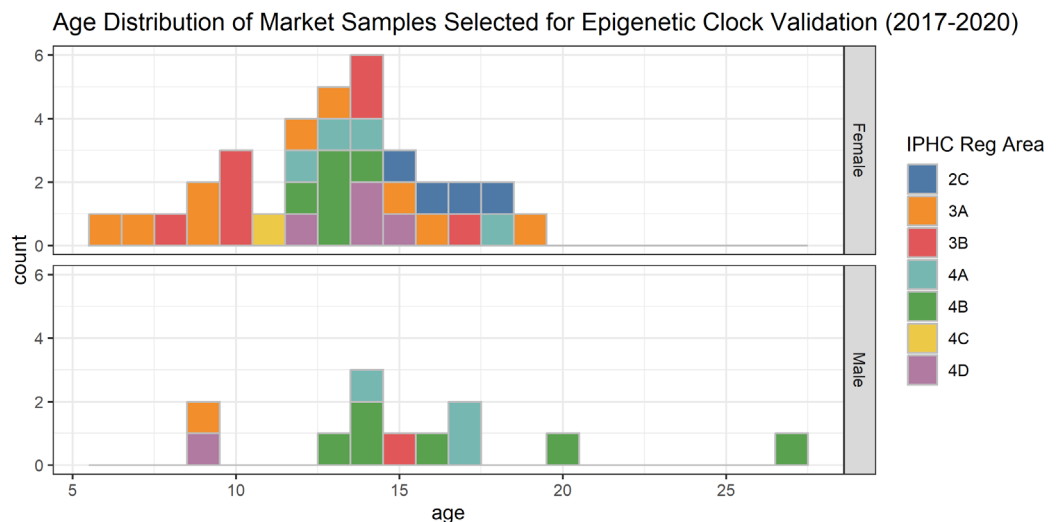


Figure 3. Age distribution of genetic samples from commercial landings (“market samples”) selected for out-of-sample validation of the Pacific halibut epigenetic clock.

- 1.2.1.3. Development of a bioinformatic workflow. A bioinformatic workflow was developed to process reduced representation bisulfite sequencing (RRBS) data in house (Figure 4). Prior to start processing the raw sequence data, a reference genome was prepared for bisulfite alignments with the Pacific halibut reference genome ([Jasonowicz et al., 2022](#); [GCF_022539355.2](#)) and control methylated and unmethylated sequences (see below) using the software Bismark.

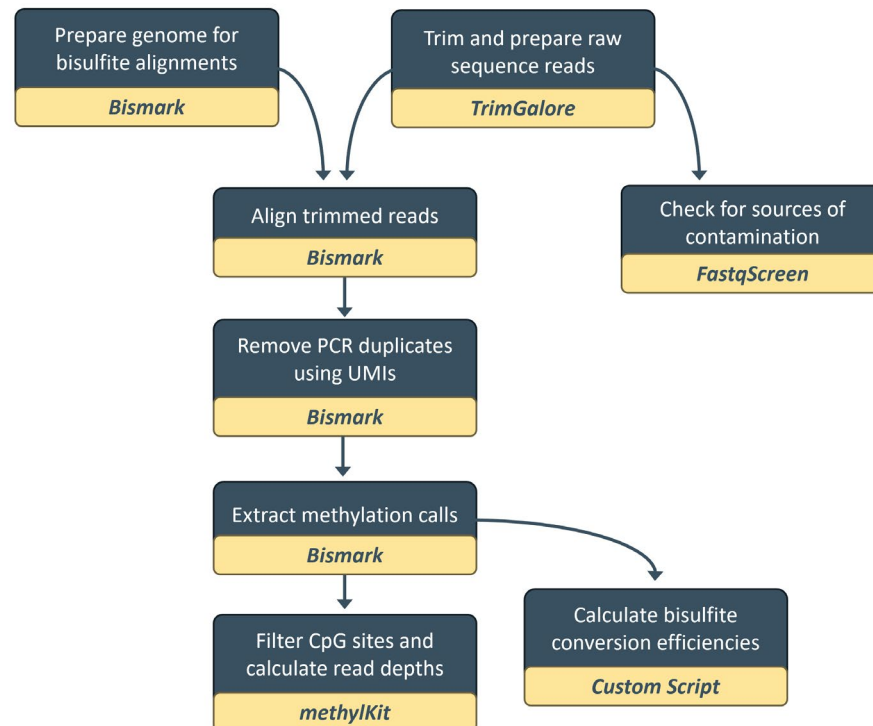


Figure 4. Basic bioinformatic workflow for processing RRBS data. The blue boxes describe each step and the software used is indicated in the yellow boxes.

1.2.1.4. Reduced representation bisulfite sequencing: library preparation and sequencing.

High-quality genomic DNA was extracted from individual fin clips and purified using the DNeasy Blood and Tissue Kit (Qiagen). Genomic DNA was used to construct individual RRBS libraries using the Premium RRBS Kit V2 (Diagenode) following the manufacturer's specifications. In brief, RRBS libraries were prepared by digesting genomic DNA with the methylation sensitive restriction enzyme *MspI*, and the resulting DNA fragments were treated with bisulfite to convert non-methylated cytosines into uracils through chemical deamination, leaving methylated cytosines unaffected. A subsequent PCR amplification step converted uracils into thymines. Constructed RRBS libraries were first assessed for concentration and fragment size distribution by TapeStation (Agilent). Subsequently, individual libraries used for the development of the epigenetic clock were pooled and sequenced on an Illumina NovaSeq-X sequencing platform in 1.5B flow cells at 2 x 50PE at the Functional Genomics Facility at the University of Chicago (Chicago, IL). In addition, individual libraries used for the out-of-sample validation of the epigenetic clock were pooled and sequenced on an Element Aviti sequencing platform at 2 x 75PE at the Functional Genomics Facility at the University of Chicago (Chicago, IL).

1.2.1.5. Sequencing data analysis and methylation calling. Prior to analysis, raw sequence reads were quality checked using *FastQC* (Andrews et al., 2015) to ensure consistent quality across sequencing runs and to identify samples that may not be suitable for

further analysis. Specifically, the raw base quality scores for each sample were used to identify samples that were poorly sequenced and should be omitted from downstream analyses. The raw sequence reads were then processed to remove Illumina adapter sequences and low quality reads using *Trim Galore!* (<https://github.com/FelixKrueger/TrimGalore>), a trimming tool designed specifically for RRBS data. Trimmed sequence reads were aligned to a bisulfite converted index of the Pacific halibut reference genome (RefSeq assembly accession: [GCF_022539355.2](https://www.ncbi.nlm.nih.gov/assembly/GCF_022539355.2)) excluding the sex chromosome (Chr09; Jasonowicz et al., 2022) to discard possible sex-associated methylation signals, using *bismark* (Krueger and Andrews, 2011) allowing for one mismatch. Having a high-quality reference genome available for Pacific halibut is a major benefit to this study as constructing one is costly and time consuming. Furthermore, the Pacific halibut genome has been annotated so that the locations and identity of genes are known, enabling the functional significance of methylated CpG sites present in protein coding gene regions to be inferred. Following alignment, polymerase chain reaction (PCR) duplicates were removed using unique molecular index (UMI) information using by supplying the `--barcode --bclconvert` options to *bismark's deduplicate_bismark* utility. Lastly, methylation levels were extracted using the *bismark_methylation_extractor* utility provided with *bismark*. For all identified CpG sites, percentage methylation was calculated as the percentage of the number of methylated reads over the number of total reads. Typically, RRBS produces in the order of hundreds of thousands of CpGs (Anastasiadi and Piferrer, 2023). Methylation levels at CpG sites for each sample were processed using the *methyKit* package (v1.24.0) (Akalın et al. 2012) in R (v4.2.3) (R Core Team 2023). To remove CpG sites covered by too few or too many reads, we retained CpG sites that were covered by at least 10 reads and did not exceed the 99.9th percentile of coverage distribution for each sample. Next, we selected CpG sites that did not have any missing data across all the samples. For each aged individual, we calculated levels of global methylation as the mean methylation percentage across the shared CpG sites identified and summarized these values for each age class using a boxplot.

- 1.2.1.6. Development of an age predicting model for Pacific halibut. The sequenced genetic samples will be randomly assigned to a training (200 samples) or a testing data set (50 samples) following an 80/20 data split. Sample assignments will be conducted using *caret* to maintain equal sex ratios in each data set. The training set will be used to fit the model and the testing set will be an independent set of data that will be used to evaluate the model fit.

The relationship between otolith-derived age and percent methylation across age-correlated CpG sites in the training data set will be characterized by performing elastic net penalized regression analysis using the R package *glmnet* (Friedman et al., 2010) set to a 10-fold cross validation with an α -parameter of 0.5 and automatically selecting the optimal penalty parameter (λ). We expect that the age-predicting model will retain in the order of a few hundred CpG sites with a low λ value. The performance of the model in the training and testing data set will be evaluated using Pearson correlations (i.e. measuring the degree of correlation between chronological and estimated age) as a measure of accuracy, MAE as a measure of precision (i.e., how well the model

fits the actual data), and relative error rates (Piferrer and Anastasiadi, 2023). Comparison of MAE between the training and testing data sets will inform on the potential overfit of the model constructed using the training data set. The linear relationship between predicted and chronological (i.e., otolith-derived) age will be visually represented and additional patterns in the data will be visualized using principal component analysis (PCA).

- 1.2.1.7. Identification of the genomic location of age markers. The Pacific halibut genome annotation (NCBI link) will also be used to determine if any functional genes are located within 400 bp of model selected CpG sites. This will inform whether clock CpG sites are proximal to specific annotated genes and whether methylation at those particular sites could have functional significance.

1.2.2. Results.

All aged fin clips used for the construction of the epigenetic clock have been processed for DNA extraction. Genomic DNA was quantified, and all samples yielded enough high-quality genomic DNA to proceed with individual library construction. Library preparation was successfully completed for all 249 individual aged samples. RRBS libraries were combined into 6 pools of 41-42 libraries each and sequenced.

Based on initial processing of the six sequenced library pools, the average PCR duplication percentage was 43.08% across all pools, ranging from 33.77% to 64.82%. The average number of deduplicated reads per individual sample ranged from 5,288,760 reads to 7,264,685 reads. Average methylated controls were 1.58% across all libraries, ranging from 1.40% to 1.86%. Average methylated levels in the control samples below 2% indicate minimal false detection of unmethylated cytosines. On the other hand, average unmethylated controls were 99.25% across all libraries, ranging from 98.95% to 99.44%, providing a positive control to assess the efficiency of bisulfite conversion of DNA. The total number of obtained deduplicated reads that were used for downstream analyses was 1,590,925,852 (i.e. over 1.5 billion reads) (Table 3).

Table 3. Initial metrics for all six library pools composed of individual libraries from project samples that will be used to develop an epigenetic clock.

Pool	Number of samples processed	Average PCR duplication rate (%)	Average number of remaining reads/sample	Average of methylated control (%)	Average of unmethylated control (%)	Sum of deduplicated reads (reads remaining)
1	42	64.82	5,288,760	1.40	98.95	222,127,927
2	41	49.56	6,506,924	1.65	99.20	266,783,900
3	42	34.99	7,264,685	1.64	99.37	305,116,779
4	42	38.35	6,509,939	1.86	99.44	273,417,435

5	41	33.77	6,443,062	1.41	99.29	270,608,611
6	41	36.99	6,167,590	1.51	99.25	252,871,200
Total	249	43.08	-	1.58	99.25	1,590,925,852

Current work is focused on processing the sequence data through the bioinformatic workflow depicted in Figure 4, with a specific focus on evaluating CpG site filtering choices that will determine which CpGs are available for the modelling step. Our current filtering process yielded 2,823,114 CpG sites per sample on average and 1,055,430 CpGs per sample on average after additional filtering based on sequence read depth (Figure 5).

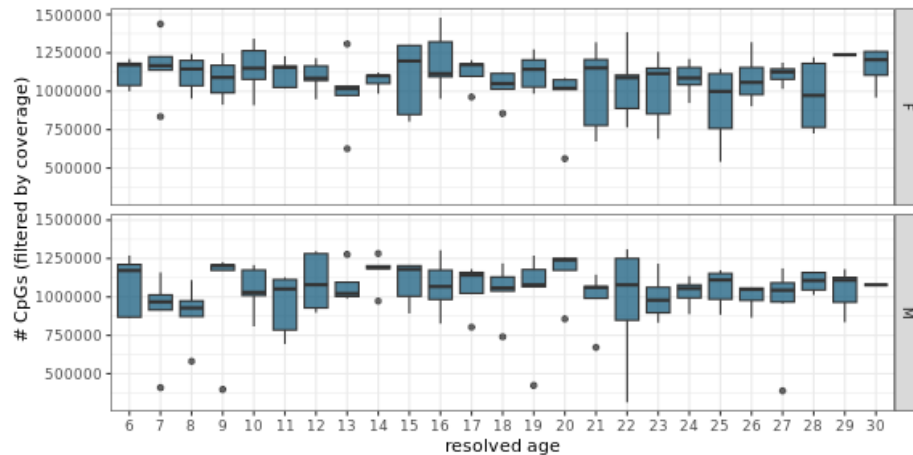


Figure 5. Number of individual CpG methylation sites by age class after filtering for sequence read depth (> 10x). Boxplots show the median (black line), 25th and 75th percentiles (box limits) and whiskers that extend to the most extreme value within 1.5x of the interquartile range.

Applying a conservative filtering approach that required complete data of all samples, 39,461 CpG sites were identified that are common to all samples. We observed a mean global methylation level of 24.6% across all aged samples (Figure 6).

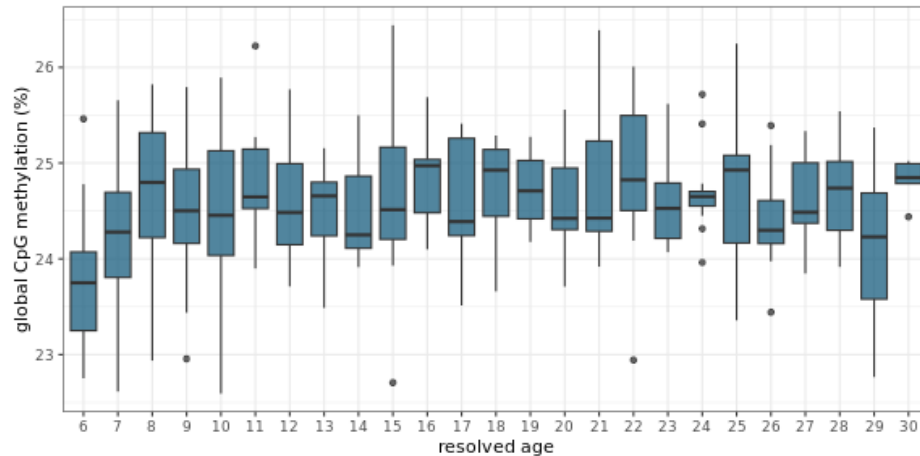


Figure 6. Individual average CpG methylation levels (%) by age class for 39,461 common CpG sites without any missing data. Boxplots show the median (black line), 25th and 75th percentiles (box limits) and whiskers that extend to the most extreme value within 1.5x of the interquartile range.

Raw sequencing data from the 46 individual libraries included in pool 7 that will be used for the out-of-sample validation of the epigenetic clock has been processed (Table 4). The average number of deduplicated reads and the total number of deduplicated reads that will be used for downstream analysis were acceptable and slightly lower than those from sequenced pools from project samples (Table 3). Bisulfate conversion efficiency was high (99.3%) and comparable to other sequenced pools.

Table 4. Initial metrics for the library pool composed of individual market samples that will be used for out-of-sample validation of the epigenetic clock.

Pool	Number of samples processed	Average PCR duplication rate (%)	Average number of remaining reads/sample	Average of methylated control (%)	Average of unmethylated control (%)	Sum of deduplicated reads (reads remaining)
7	46	28.13	4,087,418	1.74	99.27	183,933,800

In response to SRB028–Req.03 ([para. 18](#)), the IPHC Secretariat has created a preliminary full factorial design that could be used to determine whether any training-test combinations of geographic area and year perform inadequately (Figure 7).

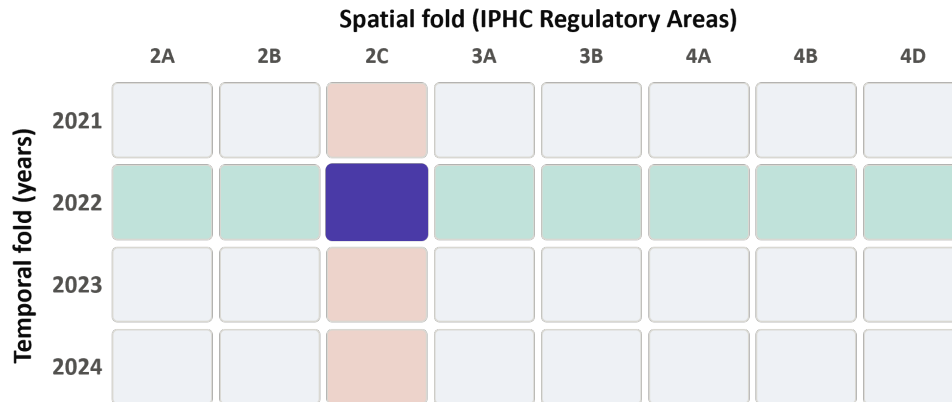


Figure 7. IPHC Regulatory Area x year holdout matrix. In this example, the model would be trained using data from every IPHC Regulatory Area and year except a specific area-year combination (blue cell: IPHC Regulatory Area 2 x Year 2022) and subsequently tested using that specific area-year combination. Alternatively, leave-on-area-out and leave-one-year-out approaches could be used to train the model with all areas (green row) or years (red column) except one (blue cell: 2C or 2022, respectively) for out-of-area and out-of-year cross validations.

In response to *SRB028–Rec.02* ([para. 35](#)) regarding the different aging methods that the IPHC Secretariat is currently using and developing, the IPHC Secretariat has compiled information on the epigenetic aging method in relation to anticipated use cases, relative development and production costs, benefits, timelines for development and decision point in Table 5.

Table 5. Characteristics of the epigenetic aging method.

Epigenetic aging	
Anticipated use cases	Additional source of ages for stock assessment Can provide insights into the biological basis of aging Could reduce uncertainty in natural mortality estimations in stock assessment
Relative costs	
Development costs	Sample processing (\$1,859) Library construction (\$17,450) Genome sequencing (\$6,000) Cloud computing services (\$3,603)
Production costs	GT-seq panel on subset of age-associated CpG sites (\$1-2 per sample)
Benefits	Non-lethal aging Useful for species with hard structures that are hard to read Accurate alternate aging method Genetic samples (fin clips) are easy to collect and store Laboratory based method Low cost, high throughput production aging

Timelines for development	Epigenetic clock: 18 months Production panel: 6 months
Decision points	Evaluation of model performance in development of the epigenetic clock Evaluation of sex, maturity status and geographic origin on age predictive capabilities Evaluation of accuracy and precision Comparison with other aging methods

2. Reproduction.

Research activities in this Research Area aim at providing information on key biological processes related to reproduction in Pacific halibut (maturity and fecundity) and to provide sex ratio information of Pacific halibut commercial landings. The relevance of research outcomes from these activities for stock assessment (SA) is in the scaling of Pacific halibut biomass and in the estimation of reference points and fishing intensity. These research outputs will result in a revision of current maturity schedules and will be included as inputs into the SA (Appendix II), and represent some of the most important biological inputs for stock assessment (please see document IPHC-2021-SRB018-06). The relevance of these research outcomes for the management and strategy evaluation (MSE) process is in the improvement of the simulation of spawning biomass in the Operating Model (Appendix III).

- 2.1. Sex ratio of the commercial landings. The IPHC Secretariat is finalizing the processing of genetic samples from the 2025 aged commercial landings.
- 2.2. Reproductive assessment. Recent sensitivity analyses have shown the importance of changes in spawning output due to changes in maturity schedules and/or skip spawning and fecundity for SA (Stewart and Hicks, 2018). Information on these key reproductive parameters provides direct input to the SA. For example, information on fecundity-at-age and -size could be used to replace spawning biomass with egg output as the metric of reproductive capability in the SA and management reference points. This information highlights the need for a better understanding of factors influencing reproductive biology and success of Pacific halibut. To fill existing knowledge gaps related to the reproductive biology of female Pacific halibut, research efforts are devoted to characterizing female reproduction in this species. Specific objectives of current studies include: 1) update of maturity schedules based on histology and calibrated visual data; and 2) fecundity estimations using the auto-diametric method.

2.2.1. Update of maturity schedules based on histology and calibrated visual data. The IPHC Secretariat provided an update on spatial and temporal patterns in maturity ogives by Biological Region from 2022 to 2024, and a revised coastwide maturity ogive using histological based data in IPHC-2025-SRB026-06. At present, the IPHC Secretariat is preparing a manuscript for publication in a peer-reviewed journal describing the temporal and spatial changes in histology-derived maturity ogives and ovarian developmental stages. Furthermore, the IPHC Secretariat developed a calibration between histological and visual maturity ogives from the 2022-2024 data and produced calibrated maturity ogives based on FISS visual maturity data from 2002-2024. During this 23-year period, we observe significant shifts in the age at 50% (A_{50}) maturity based on the calibrated maturity ogives (Figure 8; solid line). These results evidence two temporal shifts, one characterized by a gradual increase in the A_{50} (i.e. females maturing at a later age) from 2004 to 2016, and a second one characterized by a sharp decrease in A_{50} from 2017 to 2022 (i.e. females rapidly maturing at an earlier age). Studies are planned to identify possible drivers of these temporal shifts in maturity-at-age in female Pacific halibut.

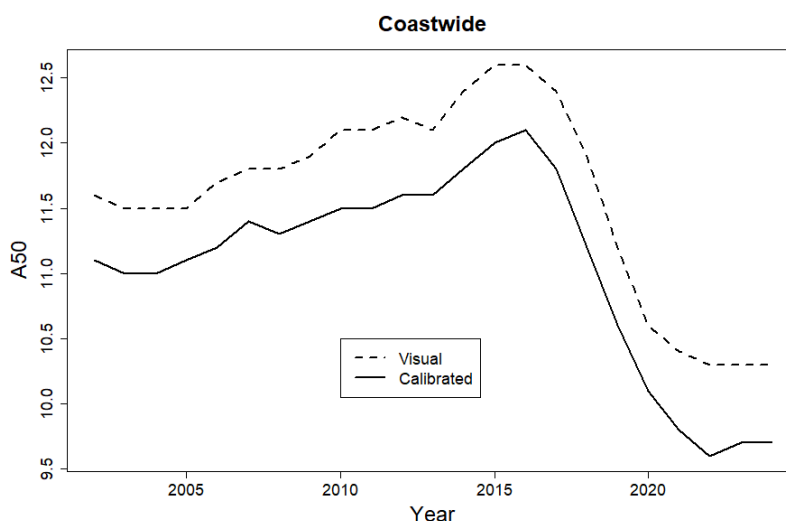


Figure 8. Coastwide A_{50} calculated from visual and calibrated maturity ogives from 2002 to 2024.

The IPHC Secretariat collected ovarian samples for maturity in the 2025 FISS, resulting in 1,276 ovarian samples collected from all four Biological Regions: 275 samples in Biological Region 2, 380 samples in Biological Region 3, 355 samples in Biological Region 4, and 266 samples in Biological Region 4B. The IPHC Secretariat is currently processing the 2025 histology samples and will present preliminary spatial and temporal analyses at SRB029. In 2026, the IPHC Secretariat will continue to collect maturity samples coastwide by targeting the collection of 400 ovarian samples in Biological Region 2, 600 samples in Biological Region 3, and 400 samples in Biological Region 4 (Figure 9). Unfortunately, no samples will be collected in Biological Region 4B in 2026 due to the lack of FISS vessel bids.

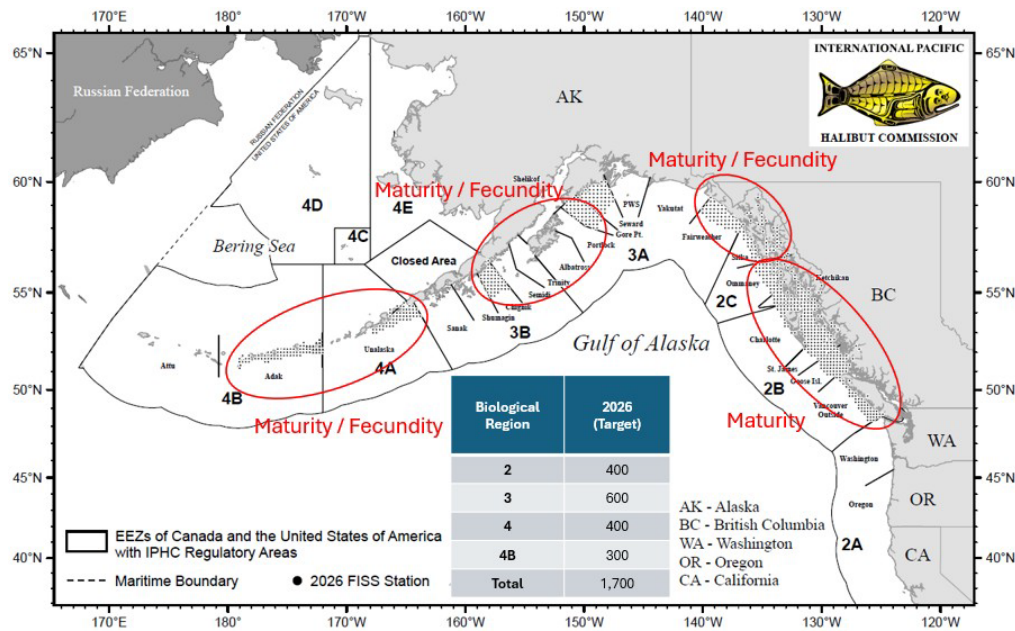


Figure 9. Coastwide map of sample areas and targets for maturity and fecundity collection in 2026 FISS.

2.2.2. Fecundity estimates. The IPHC Secretariat has initiated studies that are aimed at improving our understanding of Pacific halibut fecundity. These studies will allow us to estimate fecundity-at-size and -age and could be used to replace spawning biomass with egg output as the metric for reproductive capability in SA and management reference points. Fecundity determinations will be conducted using the auto-diametric method (Thorsen and Kjesbu, 2001; Witthames et al., 2009) and IPHC Secretariat staff received training on this method by experts in the field (NOAA Fisheries, Northeast Fisheries Science Center, Wood Hole, MA) in May 2023. Ovarian samples for the development and application of the auto-diametric method to estimate fecundity in female Pacific halibut have been collected during the FISS in 2023, 2024 and 2025, as well as two special collections in IPHC Regulatory Area 2B in 2024 and 2025 (Figure 10). In 2023, sampling was conducted only in Biological Region 3, with a total of 452 fecundity samples collected. In 2024, sampling was conducted in Biological Regions 2 and 4, with 149 and 359 fecundity samples collected, respectively. In the Fall (Oct/Nov) of 2024, 271 additional fecundity samples targeting large females (85-200+ cm in fork length) were collected in Biological Region 2. This sampling was conducted to collect later developing females to help build the auto-diametric curve for fecundity estimations. For 2025, in addition to 878 samples collected in all four Biological Regions in the FISS, 242 fecundity samples were collected in Biological Region 2 in a special project targeting large females during the late summer months. This comprehensive collection of ovarian samples will be used initially for the development of the auto-diametric method, followed by actual fecundity estimations by age and by size (length and weight).

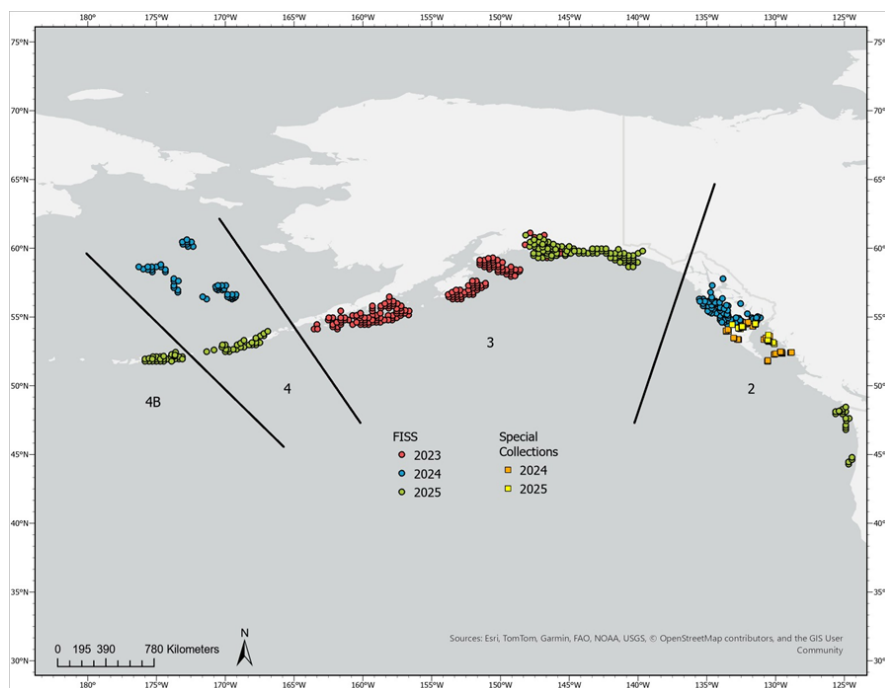


Figure 10. Coastwide map of 2023, 2024 and 2025 samples for fecundity collected in FISS (circle colors), and 2024 and 2025 special collection fecundity samples in IPHC Regulatory Area 2B (square colors).

In 2026, the IPHC Secretariat will continue to collect ovarian samples during the FISS for fecundity estimation. Specifically, whole ovary weights and samples are being collected from G2 (i.e. mature) females in Biological Region 2 (Sitka charter region), Biological Region 3 (Fairweather, Gore Pt., and Chignik charter regions), and Biological Region 4 (Unalaska charter region). The IPHC Secretariat will also be collecting fecundity samples from mid-September to October of 2026 (outside of FISS) during a special project in Biological Region 2 (Ommaney charter region) and Biological Region 3 (Gore Pt. charter region). The target sample collection for each Biological Region is 150 samples of >90cm G2 females. These samples will allow the IPHC Secretariat to spatially compare fecundity estimations among multiple Biological Regions, as previous special project collections (2024 and 2025) were conducted in Biological Region 2.

From 2023-2025, a total of 1,154 G2 ovarian samples have been collected for fecundity estimations. When examining the age and weight distribution of mature (G2) females collected for fecundity sampling (Figures 11-12), the distribution of fish appeared to be skewed to the right for both parameters, but more pronounced for weight (Figure 12). For the mature female samples collected from 2023-2025, the total range of ages was from 6 to 38 years old, and the total range of weights was from 2 to 92 kg. The largest proportion of sampled fish was from 11 to 13 years old, and from 3 to 9 kg in weight.

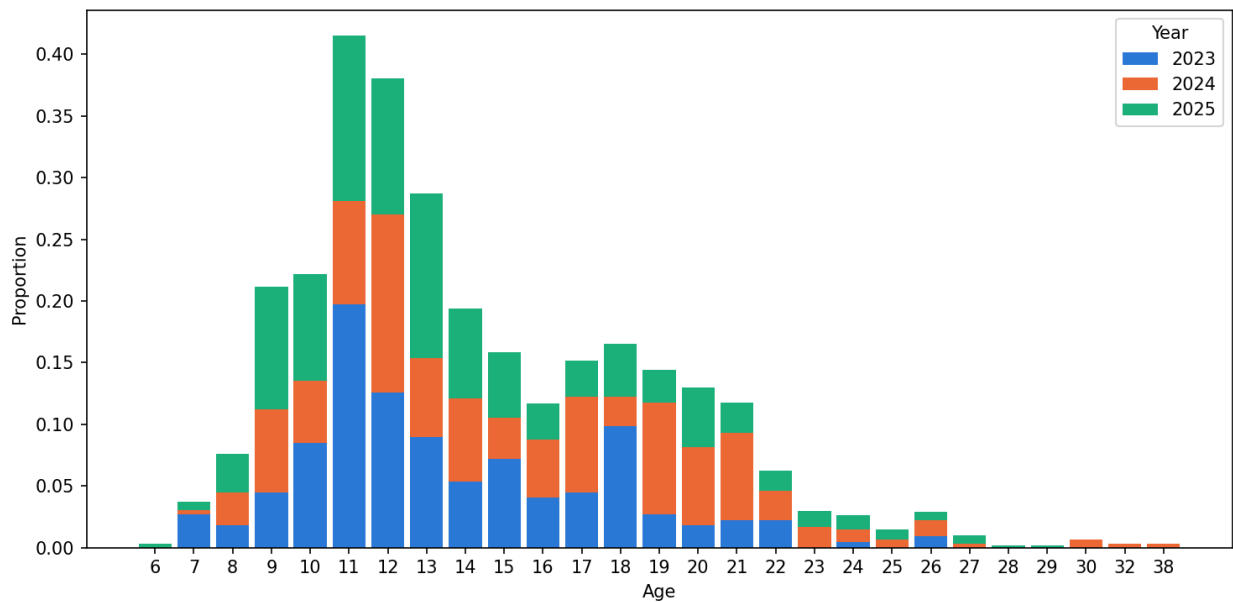


Figure 11. Stacked bar chart showing proportions of age (years) of mature (G2) female Pacific halibut collected for fecundity estimation in 2023, 2024, and 2025.

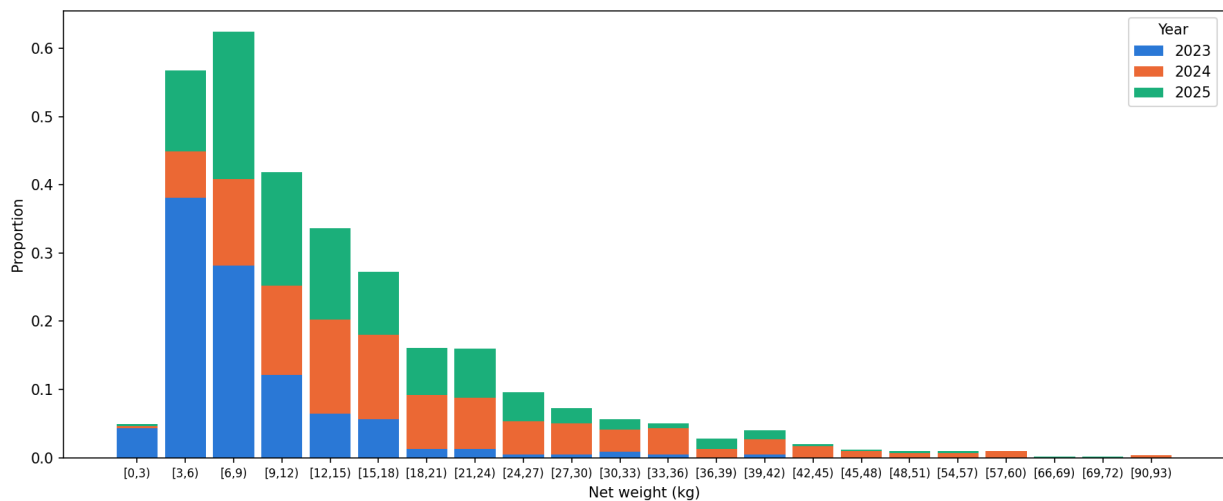


Figure 12. Stacked bar chart showing proportions of net weight (kg) of mature (G2) female Pacific halibut collected for fecundity estimation in 2023, 2024, and 2025.

3. Growth.

Research activities conducted in this Research Area aim at providing information on somatic growth processes driving size-at-age in Pacific halibut. The relevance of research outcomes from these activities for stock assessment (SA) resides, first, in their ability to inform yield-per-recruit and other spatial evaluations for productivity that support mortality limit-setting, and, second, in that they may provide covariates for projecting short-term size-at-age and

may help delineate between fishery and environmental effects, thereby informing appropriate management responses (Appendix II). The relevance of these research outcomes for the management and strategy evaluation (MSE) process is in the improvement of the simulation of variability and to allow for scenarios investigating climate change (Appendix III).

The IPHC Secretariat has conducted studies aimed at elucidating the drivers of somatic growth leading to the decline in SAA by investigating the physiological mechanisms that contribute to growth changes in the Pacific halibut. The two main objectives of these studies have been: 1) the identification and validation of physiological markers for somatic growth; and 2) the application of molecular growth markers for evaluating growth patterns in the Pacific halibut population. A manuscript describing the results of these studies has been published (Planas et al., 2025).

4. Mortality and Survival Assessment.

Information on all Pacific halibut removals is integrated by the IPHC Secretariat, providing annual estimates of total mortality from all sources for its stock assessment. Bycatch and wastage of Pacific halibut, as defined by the incidental catch of fish in non-target fisheries and by the mortality that occurs in the directed fishery (i.e. fish discarded for sublegal size or regulatory reasons), respectively, constitute important sources of mortality that can result in significant reductions in exploitable yield in the directed fishery. Given that the incidental mortality from the commercial Pacific halibut fisheries and bycatch fisheries is included as part of the total removals that are accounted for in stock assessment, changes in the estimates of incidental mortality will influence the output of the stock assessment and, consequently, the catch levels of the directed fishery. Research activities conducted in this Research Area aim at providing information on discard mortality rates and producing guidelines for reducing discard mortality in Pacific halibut in the longline and recreational fisheries. The relevance of research outcomes from these activities for stock assessment (SA) resides in their ability to improve trends in unobserved mortality to improve estimates of stock productivity and represent the most important inputs in fishery yield for stock assessment (Appendix II). The relevance of these research outcomes for the management and strategy evaluation (MSE) process is in fishery parametrization (Appendix III).

- 4.1. Estimation of discard mortality rates in the charter recreational sector. Results from a recently completed study investigating discard mortality rates and characteristics of fish captured and released using guided recreational fishery practices are currently being prepared for publication in a peer-reviewed journal.

5. Fishing technology.

The IPHC Secretariat has determined that research to provide the Pacific halibut fishery with tools to reduce whale depredation is considered a high priority (Appendix I). This research is now contemplated as one of the research areas of high priority within the 5-year Program of Integrated Research and Monitoring (2022-2026). Towards this goal, the IPHC secretariat has been investigating gear-based approaches to catch protection as a means for minimizing

whale depredation in the Pacific halibut and other longline fisheries with funding from NOAA's Bycatch Research and Engineering Program (BREP) (NOAA Awards NA21NMF4720534 and NA23NMF4720414; Appendix IV). The results and outcomes of the initial pilot phase of this project were reported in the documentation provided for the previous SRB meetings: IPHC-2022-SRB020-08 and IPHC-2024-SRB024-09.

The second phase of this project focused on further refinement and performance characterization of the shuttle device (Figure 13A) in the presence of toothed whales in IPHC Regulatory Area 4A. Field operations occurred from 21-28 May 2025 aboard the F/V Oracle (17.5 m, 58 ft) in the Bering Sea and Aleutian Islands off Alaska in known depredation hotspots and with higher expected Pacific halibut catch rates than seen in the pilot testing.

Eighteen sets were successfully completed, generating 15 sets of shuttle and control catch comparison data. Depredating orcas (identified by eaten or damaged Pacific halibut and by orca presence (Figure 13B,C) were present at 6 of the paired sets. Camera systems developed by project participants were successfully deployed both on the control gear and on the shuttle itself (exterior and interior forward and reverse views), capturing approximately 80 hours of underwater footage combined, enabling us to better quantify shuttle performance and retention rates (Figure 13D). Approximately 73 hours (11/15 paired sets) of the footage has been reviewed in detail, and 5,463 hook status observations have been recorded across all four cameras.

Preliminary results from video footage suggest that the shuttle device had high retention rates of Pacific halibut when compared to other frequently encountered species (Table 6). In addition, preliminary results on catch rate comparisons between protected and unprotected skates did not show clear differences, likely due to high variability among catch rates within treatments. Overall, the shuttle device demonstrated good retention capacity (up to 835 kg/1,800 lb) in the trials, with capacity for higher weights. Incidentally, rare footage of Orcas around the groundline at depth was captured.

Table 6. Numbers of fish encountered by the shuttle device that were either excluded, entered and escaped, entered and passed through still on the hook, and/or finally retained on 11 of 15 sets with video footage analyzed to date.

Common Name	Encountered	Excluded	Entered and Escaped	Entered and Passed Through	Retained
Pacific halibut	121	1 (0.8%)	0	9 (7.5%)	112 (93.3%)
Sablefish	186	2 (1.1%)	54 (29.4%)	34 (18.5%)	96 (52.2%)
Pacific cod	124	3 (2.4%)	13 (10.7%)	6 (5.0%)	102 (84.3%)
Rockfish	19	7 (43.8%)	2 (16.7%)	1 (8.3%)	9 (75.0%)
Skate	18	3 (16.7%)	0	2 (13.3%)	13 (86.7%)

Data analysis and video review are ongoing. These preliminary results suggest potential for this approach to interrupt the reward cycle underpinning toothed whale depredation.

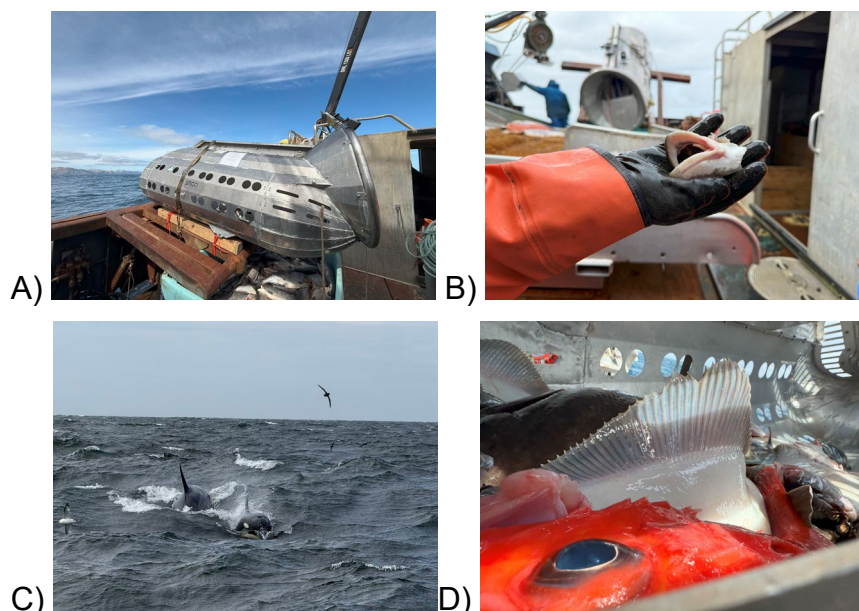


Figure 13. A) Shuttle device in transport. B) Typical evidence (lips only) of depredation. C) Killer whales rapidly approaching the hauling site. D). Catch retained within the shuttle.

RECOMMENDATION/S

That the SRB:

- a) **NOTE** paper IPHC-2026-SRB029-05 which provides a response to Requests and Recommendations from SRB028, and a report on current biological research activities contemplated within the IPHC's five-year Program of Integrated Research and Monitoring (2022-26).

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APPENDIX I

Integration of biological research, stock assessment (SA) and management strategy evaluation (MSE): rationale for biological research prioritization

Research areas	Research activities	Research outcomes	Relevance for stock assessment	Relevance for MSE	Specific analysis input	SA Rank	MSE Rank	Research prioritization
Migration and population dynamics	Population structure	Population structure in the Convention Area	Altered structure of future stock assessments	Improve parametrization of the Operating Model	If 4B is found to be functionally isolated, a separate assessment may be constructed for that IPHC Regulatory Area	2. Biological input	1. Biological parameterization and validation of movement estimates and recruitment distribution	2
	Distribution	Assignment of individuals to source populations and assessment of distribution changes	Improve estimates of productivity		Will be used to define management targets for minimum spawning biomass by Biological Region	3. Biological input		2
	Larval and juvenile connectivity studies	Improved understanding of larval and juvenile distribution	Improve estimates of productivity		Will be used to generate potential recruitment covariates and to inform minimum spawning biomass targets by Biological Region	3. Biological input	1. Biological parameterization and validation of movement estimates	2
Reproduction	Histological maturity assessment	Updated maturity schedule	Scale biomass and reference point estimates	Improve simulation of spawning biomass in the Operating Model	Will be included in the stock assessment, replacing the current schedule last updated in 2006	1. Biological input		1
	Examination of potential skip spawning	Incidence of skip spawning			Will be used to adjust the asymptote of the maturity schedule, if/when a time-series is available this will be used as a direct input to the stock assessment			1
	Fecundity assessment	Fecundity-at-age and -size information			Will be used to move from spawning biomass to egg-output as the metric of reproductive capability in the stock assessment and management reference points			1
	Examination of accuracy of current field macroscopic maturity classification	Revised field maturity classification			Revised time-series of historical (and future) maturity for input to the stock assessment			1
Growth	Evaluation of somatic growth variation as a driver for changes in size-at-age	Identification and application of markers for growth pattern evaluation	Scale stock productivity and reference point estimates	Improve simulation of variability and allow for scenarios investigating climate change	May inform yield-per-recruit and other spatial evaluations of productivity that support mortality limit-setting		3. Biological parameterization and validation for growth projections	5
		Environmental influences on growth patterns			May provide covariates for projecting short-term size-at-age. May help to delineate between effects due to fishing and those due to environment, thereby informing appropriate management response			5
		Dietary influences on growth patterns and physiological condition			May provide covariates for projecting short-term size-at-age. May help to delineate between effects due to fishing and those due to environment, thereby informing appropriate management response			5
Mortality and survival assessment	Discard mortality rate estimate: longline fishery	Experimentally-derived DMR	Improve trends in unobserved mortality	Improve estimates of stock productivity	Will improve estimates of discard mortality, reducing potential bias in stock assessment results and management of mortality limits	1. Fishery yield	1. Fishery parameterization	4
	Discard mortality rate estimate: recreational fishery				Will improve estimates of discard mortality, reducing potential bias in stock assessment results and management of mortality limits			4
	Best handling and release practices	Guidelines for reducing discard mortality			May reduce discard mortality, thereby increasing available yield for directed fisheries	2. Fishery yield		4
Fishing technology	Whale depredation accounting and tools for avoidance	New tools for fishery avoidance/deterrence; improved estimation of depredation mortality	Improve mortality accounting	Improve estimates of stock productivity	May reduce depredation mortality, thereby increasing available yield for directed fisheries. May also be included as another explicit source of mortality in the stock assessment and mortality limit setting process depending on the estimated magnitude	1. Assessment data collection and processing		3



APPENDIX II

List of ranked biological uncertainties and parameters for stock assessment (SA) and their links to biological research areas and research activities

SA Rank	Research outcomes	Relevance for stock assessment	Specific analysis input	Research Area	Research activities
1. Biological input	Updated maturity schedule	Scale biomass and reference point estimates	Will be included in the stock assessment, replacing the current schedule last updated in 2006	Reproduction	Histological maturity assessment
	Incidence of skip spawning		Will be used to adjust the asymptote of the maturity schedule, if/when a time-series is available this will be used as a direct input to the stock assessment		Examination of potential skip spawning
	Fecundity-at-age and -size information		Will be used to move from spawning biomass to egg-output as the metric of reproductive capability in the stock assessment and management reference points		Fecundity assessment
	Revised field maturity classification		Revised time-series of historical (and future) maturity for input to the stock assessment		Examination of accuracy of current field macroscopic maturity classification
2. Biological input	Stock structure of IPHC Regulatory Area 4B relative to the rest of the Convention Area	Altered structure of future stock assessments	If 4B is found to be functionally isolated, a separate assessment may be constructed for that IPHC Regulatory Area	Genetics and Genomics	Population structure
3. Biological input	Assignment of individuals to source populations and assessment of distribution changes	Improve estimates of productivity	Will be used to define management targets for minimum spawning biomass by Biological Region		Distribution
	Improved understanding of larval and juvenile distribution		Will be used to generate potential recruitment covariates and to inform minimum spawning biomass targets by Biological Region	Migration	Larval and juvenile connectivity studies
1. Assessment data collection and processing	Sex ratio-at-age	Scale biomass and fishing intensity	Annual sex-ratio at age for the commercial fishery fit by the stock assessment	Reproduction	Sex ratio of current commercial landings
	Historical sex ratio-at-age		Annual sex-ratio at age for the commercial fishery fit by the stock assessment		Historical sex ratios based on archived otolith DNA analyses
2. Assessment data collection and processing	New tools for fishery avoidance/deterrence; improved estimation of depredation mortality	Improve mortality accounting	May reduce depredation mortality, thereby increasing available yield for directed fisheries. May also be included as another explicit source of mortality in the stock assessment and mortality limit setting process depending on the estimated magnitude	Mortality and survival assessment	Whale depredation accounting and tools for avoidance
1. Fishery yield	Physiological and behavioral responses to fishing gear	Reduce incidental mortality	May increase yield available to directed fisheries	Mortality and survival assessment	Biological interactions with fishing gear
2. Fishery yield	Guidelines for reducing discard mortality	Improve estimates of unobserved mortality	May reduce discard mortality, thereby increasing available yield for directed fisheries	Mortality and survival assessment	Best handling practices: recreational fishery

APPENDIX III

List of ranked biological uncertainties and parameters for management strategy evaluation (MSE) and their links to biological research areas and research activities

MSE Rank	Research outcomes	Relevance for MSE	Research Area	Research activities
1. Biological parameterization and validation of movement estimates	Improved understanding of larval and juvenile distribution	Improve parametrization of the Operating Model	Migration	Larval and juvenile connectivity studies
	Stock structure of IPHC Regulatory Area 4B relative to the rest of the Convention Area			Population structure
2. Biological parameterization and validation of recruitment variability and distribution	Assignment of individuals to source populations and assessment of distribution changes	Improve simulation of recruitment variability and parametrization of recruitment distribution in the Operating Model	Genetics and Genomics	Distribution
	Establishment of temporal and spatial maturity and spawning patterns	Improve simulation of recruitment variability and parametrization of recruitment distribution in the Operating Model	Reproduction	Recruitment strength and variability
3. Biological parameterization and validation for growth projections	Identification and application of markers for growth pattern evaluation	Improve simulation of variability and allow for scenarios investigating climate change	Growth	Evaluation of somatic growth variation as a driver for changes in size-at-age
	Environmental influences on growth patterns			
	Dietary influences on growth patterns and physiological condition			
1. Fishery parameterization	Experimentally-derived DMRs	Improve estimates of stock productivity	Mortality and survival assessment	Discard mortality rate estimate: recreational fishery



APPENDIX IV
Summary of current external research grants

Project #	Grant agency	Project name	PI	Partners	IPHC Budget (\$US)	Management implications	Grant period
1	Alaska Sea Grant	Development of a non-lethal genetic-based method for aging Pacific halibut (R/2024-05)	IPHC, Alaska Pacific Univ. (APU)	Alaska Fisheries Science Center-NOAA (Juneau)	\$60,374	Stock structure	December 2024-January 2027
Total awarded (\$)					\$60,374		