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#### **Brief Communication**

Is bigger really better? Towards improved models for testing how Atlantic salmon Salmo salar smolt size impacts marine survival

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Running head Smolt size and marine survival

#### Abstract

A general framework is presented that should enhance our understanding of how intrinsic factors, such as body size, and extrinsic factors, such as climate, affect the dynamics and demographics of fish populations. Effects of intrinsic factors, notably studies relating juvenile Atlantic salmon *Salmo salar* body size to their probability to return as an adult, are often context-dependent and anecdotal, due to data constraints. By merit of its flexible specification, this framework should admit datasets with a range of situation-specific nuances, collected using different approaches, and thereby deliver more general and robust findings for more effective population management.

Key words: smolt body size, migration, lifecycle, state-space model, Bayesian

There are few wild populations unaffected by human-induced environmental changes, such as climate change, overexploitation, invasive species and their synergies (Brook, Sodhi & Bradshaw, 2008). For example, the abundance of Atlantic salmon *Salmo salar* (taken as their nominal catches) has declined precipitously across their range since the 1970s (ICES, 2017; Fig. 1), probably due to a range of interacting factors (Limburg & Waldman, 2009; Mills *et al.*, 2013), with populations now often augmented by hatchery-reared fish (Aprahamian *et al.*, 2003; Molony *et al.* 2003).

To manage populations effectively generally requires understandings of how intrinsic and extrinsic factors, and their interactive and legacy effects, affect individual traits and behaviours (Clutton-Brock & Sheldon, 2010). Extrinsic factors potentially implicated in S. salar population declines include climate-driven changes in sea conditions and planktonic communities (Beaugrand & Reid, 2012), predation (e.g., Riley et al., 2011) and the timing of seaward emigrating juvenile salmon (known as *smolt*) migration associated with climate (e.g., Otero et al., 2014). However, there is growing evidence that intrinsic factors carried over from their freshwater stages are important in marine mortality, such as their body size and/ or condition at smolting (Russell et al., 2012). Should the effects of their freshwater life-phase strongly influence their marine survival, then this would have fundamental implications for smolt management because it would promote strategies that maximise not just the number of smolts but also their quality (Russell et al., 2012). Furthermore, it is important to account for such legacy effects in modelling the respective contributions of different factors to overall change in population strength. Correspondingly, the aim of this study was to explore, through literature review, the potential influences of S. salar body size at smolting on their subsequent marine survival, and consider how this can be tested more robustly, for example, by accounting for imperfect detection.

At a general level, theory suggests that smolt mortality might be inversely related to their body size, i.e., the inverse-weight hypothesis (Ricker, 1976). Many studies have provided some empirical evidence testing this 'bigger is better' paradigm (sensu Sogard, 1997). Koenings et al. (1993) suggested a positive influence of smolt length on marine survival in 12 populations, although the pattern was non-linear across age groups and exacerbated by latitudinal variation. Several studies, including Henderson & Cass (1991) and Holtby et al. (1990), revealed surviving S. salar smolts were generally of greater length (as estimated by scale back-calculations) than the mean length of their corresponding cohort. While the bigger is better paradigm could reflect the consequences of general processes, such as avoiding gape-limited predators and increasing prey options, it might not be universal. For example, medium-sized smolts had the highest marine survival rates in the River Imsa, Norway, although the relatively low survival of larger smolts could not be disentangled from the influence of their emigration timing (Jonsson et al. 2017). Armstrong et al. (2017) and Saloniemi et al. (2004) both provided strong arguments that larger smolt body sizes increased marine survival, where both utilised individual-level data and considered covariates and their interactions (Fig. 2). However, when assessed across a larger number of studies assessing the influence of smolt length on subsequent marine survival, support for the bigger is better paradigm seems equivocal (Table 1).

There are some patterns evident in Table 1 that can be used for formulating future studies and model development. Most of the studies regress a time series of mean lengths on a time series of marine survival (usually expressed as "adult return rate", which measures individual probability to return as an adult irrespective of time spent at sea) for a single river or stock (Type 1 in Table 1). There are two exceptions to this approach that regress the same variables but for 6 stocks (Dempson *et al.*, 2003) and 12 stocks (Koenings *et al.*, 1993), although the latter does not account for stock in the statistical model, risking possible pseudo-

replication. Another group of studies use back-calculated lengths from scales (Type 2 in Table 1), which introduces a non-quantified uncertainty due to measurement and model choice. Another group of studies examine the fate of tracked individuals (via telemetry) and examines the influence of length class on their survival (Type 3 in Table 1). The final group of studies presents plots of patterns but with no formal statistical analyses to quantify length effects (Type 4 in Table 1). Most studies also tend to use time series data on *S. salar* and consider year as the unit of variance. This is not surprising, since most of knowledge on *S. salar* marine survival comes from long-term monitoring programmes (ICES, 2017). This is, at least in part, due to difficulties associated with studying individuals and populations at sea, although telemetry studies are now able to provide movement data from estuarine and even near-shore coastal environments (e.g., Newton *et al.*, 2016).

Statistically, only 4 of the studies considered covariates to either represent the variance fairly, i.e., to avoid pseudo-replication, or as potential competing hypotheses. In each case, the covariates were important to the study findings and thus there is a strong case for using covariates in future models. A good example is provided by Armstrong *et al.* (2017), who used individual-level covariates to generalise their findings beyond the years sampled (by using a random year effect) and to examine evidence for competing hypotheses (body condition and migration timing). Among the studies of Table 1, all but one considered linear terms only, despite acknowledging their inadequacy for some of the datasets (e.g., Holtby *et al.*, 1990).

Most of the studies in Table 1 benefitted from monitoring programmes that provided long time series and large numbers of smolts. Such large numbers, even when stratified by year, afford a good representation of the sample mean – the response variable most commonly used. However, using the individual data, where available, could provide greater insight, especially for individuals at the limits of the population length range and where using population means is not meaningful. For example, Saloniemi *et al.* (2004) used logistic regression to examine the effect of individual smolt length, relevant covariates and their interactions to reveal a positive effect of length on marine survival. Moreover, their use of individual-level data meant they required only two years of data and a moderate sample size (Table 1; Fig. 2). While a rich source of individual length data could be sourced from scale analyses and back-calculation (as per the Type 2 studies in Table 1), this requires careful consideration as: (i) lengths back-calculated from scales are subject to uncertainty in the model used and its parameters (Francis, 1990); (ii) scale collection protocol could be biased towards individuals of common characteristics, e.g., larger individuals sought by anglers; and (iii) if comparing the back-calculated lengths to the pool of observed lengths (e.g., Henderson & Cass, 1991), it should be considered that the denominator (the pool of observed lengths) might include the numerator (the back-calculated lengths).

If individual-level information is lacking from 'data-rich' long-term monitoring programmes, then an alternative might be to use abundant short and noisy datasets from 'data-poor' fisheries (Bentley, 2015). For example, Koenings *et al.* (1993) used short time series (1 to 9 years) from 12 stocks to suggest a positive effect of length on subsequent marine survival, albeit that they used annual mean data and omitted factors to allow for baseline differences between rivers and years (*c.f.* Armstrong *et al.*, 2017). Methods exist that can integrate small and noisy datasets to tease out common signals, and these methods can also admit missing data, which is often a feature of these datasets (Bentley, 2015).

As most of the datasets in Table 1 utilise mark-recapture methods, then their data also present a potential confound to marine survival estimates, as not all individuals are redetected, i.e., detection is imperfect. Detection efficiency is a measure of the probability (p)that a device (or array of devices) detects a tag moving within the area that the device was installed to monitor, which under perfect conditions will be p = 1. Many factors will cause p < 1, including animal behaviour, which might relate to size, and environmental conditions. Imperfect detection is the term used to describe the effect of these factors on perfect detection. There is a large and growing literature highlighting the importance of imperfect detection, factors affecting it and methods that can account for it (e.g., Guillera-Arroita, 2017), including a class of models that separate observation and process errors, commonly called state-space models (e.g., Gimenez et al., 2007). Failure to account for imperfect detection, particularly when the probability of detecting an individual is low to moderate (e.g., < 90%) or the sample size is low, can result in imprecise inferences that are biased and inaccurate (e.g., Gimenez et al., 2007). This is particularly concerning when interpreting data from telemetry studies that usually have low to moderate detection rates and low sample sizes. For example, Newton et al. (2016) studied the effect of tagging on survival of smolts migrating through Lough Foyle, Ireland, and found no evidence that smaller smolts were less likely to survive to be detected exiting the lough to sea, although they could not disregard the possibility that the 8 of 33 unsuccessful lough migrants (or indeed the 27 smolts not detected entering the lough) were simply not detected. Imperfect detection is likely to affect most studies in Table 1, and its effect should not be neglected.

Given the issues outlined above, it is suggested that state-space models (SSM) are well suited to future testing of the bigger is better paradigm for migrating *S. salar* smolts. These explicitly model the underlying ecological or state process (Equation 1), e.g., the effect of smolt size on its marine survival, and the observational process (Equation 2), e.g., the probability of detecting a surviving smolt. When formulated in a Bayesian language (e.g., Just Another Gibbs Sampler [JAGS]: <u>http://mcmc-jags.sourceforge.net/</u>), they amount to a set of deterministic and stochastic equations. In the simple case of estimating the effect of length on the survival probability of smolt *i* in a single river, then:

$$y_{i,t}|z_{i,t} \sim \text{Bernoulli}(z_{i,t}p)$$
 (1)

$$z_{i,t+1}|z_{i,t} \sim \text{Bernoulli}(z_{i,t}\phi_i) \tag{2}$$

where t > 0, z is a latent variable describing the state of smolt *i* at time t,  $\phi_i$  is the survival rate of smolt *i* from state  $z_{i,t}$  to state  $z_{i,t+1}$  and y is the observation of that smolt given the probability p of detecting it. From these equations, it can be noted that  $\phi_i$  and p are timeinvariant and p does not vary for individuals. To estimate the effect of smolt *i* length  $l_i$  on its survival,  $\phi_i$  is specified as a deterministic function of logistic regression parameters:

$$logit(\phi_i) = \alpha + \beta_1 l_i \tag{3}$$

where  $\alpha$  is the estimated marine survival of any smolt returning to our river and  $\beta_1$  is the effect of smolt *i* length on  $\alpha$ , while accounting for imperfect detection, i.e., 1 - p.

The ecological applications of SSM have increased due, at least in part, to their flexibility (Royle & Dorazio, 2008). For example, Gimenez et al. (2007) provide an instructive overview of SSM theory and an accompanying illustration using individual mark and recapture data collected on the European dipper Cinclus cinclus. Holbrook et al. (2014) uses SSM to estimate sea lamprey Petromyzon marinus passage through a dam using individual acoustic tagging data. A few SSM extensions are also worth noting. Equation 3 can be modified through additional covariates that are measured at the level of individual, group, or stock and are included by specifying coefficients for their (fixed) effects. For example, an effect of fat content of smolt *i* could be estimated by including the term  $\beta_2 w_i$  in Equation 3. Care should be taken to ensure the effects are indexed at the correct level. For example, a fixed effect of river is included with the term  $\beta_3 r$  that adds another stratum to all other effects, i.e., length is measured for smolts emigrating from river r in 1, 2, ..., R (where R is the number of rivers) and is therefore indexed with r as  $l_{i,r}$ . Note, by leaving  $\beta_1$  unindexed, the effect of smolt length is estimated assuming that it is identical across rivers. It is a small step to specifying river as a random effect, i.e., acknowledging differences between rivers but treating rivers as a sample of a larger "population" of rivers: rather than specifying  $\beta_3$  as a single coefficient, it is specified it as a vector of coefficients  $\beta_3$ , with effects drawn from a distribution defined by a common mean effect and variance (see Kéry & Schaub [2011] for a more complete description). With these extensions, it is straightforward to include fixed effects (e.g., latitude; Koenings *et al.*, 1993) or random effects (e.g., year; Armstrong *et al.*, 2017).

A further strength of SSM, and hierarchical models more generally, is the idea that information contained in short and noisy datasets can use information from larger, longer and less noisy datasets (Parent & Rivot, 2012). Assuming a single stock dataset per river (although this could be relaxed), then this is achieved by including a random effect of river. Although both datasets provide information to update the common mean effect estimate (and its variance), presumably the longer and less noisy dataset is providing more information, which is transferred to the shorter and noisier dataset.

There are assumptions inherent in each approach in Table 1. For example, tagging studies generally use a constant tag size, which is a higher, albeit not necessarily significant, burden on smaller fish. For example, survival effects of some tags, e.g., passive integrated transponder tags (~0.1g), are considered negligible while the effects of larger tags, e.g., acoustic telemetry transmitters (>1.0g), deserve more consideration. To test for effects of a constant size tag on variable sized smolts requires a baseline understanding of how survival relates to smolt size in untagged fish. It is not valid to infer no effect of tags from an absence of a significant size-mortality effect in a group of tagged fish alone (Newton *et al.* 2016) because there is no control to inform on how mortality would relate to size in a particular study situation in the absence of tagging. Variation in tag effect with smolt size could, however, be quantified in an SSM by, for example, contrasting Type 1 (tagging) with Type 2 studies, in which fish handling and tagging is not a consideration. In this case, the Type 2 approach would provide the control situation.

Estimating SSM parameters by Monte Carlo Markov Chains (MCMC) allows for the natural expression and propagation of uncertainties in parameter estimates to model outputs. Correctly parameterised, uncertainties from other sources could also propagate through the model. For example, mark-recapture studies rely on detection devices that can fail, which would enter the model as missing data rather than removing them. Changes to the monitoring apparatus, e.g., loss or addition of a new acoustic receiver, could be accounted for in a similar manner. Another source of uncertainty is model choice. For example, uncertainty in the model used to estimate smolt length from scales through back-calculation could be captured in a SSM, either through prior information or by implementing the back-calculation within the SSM itself. Similarly, acoustic tracking data, which provide information about estuarine and near-shore coastal mortality, could be admitted directly or indirectly. SSM that accommodate information from different data sources are commonly referred to as Integrated Population Models, and their use in ecology is increasing (e.g., Robinson *et al.*, 2014).

Although strongly advocating a move towards a general SSM to test the bigger is better paradigm for *S. salar* smolts, these models should not be considered as a panacea, as they too can have estimation problems when the process error is swamped by measurement error (Auger-Méthé *et al.*, 2016). Consequently, this study can be considered as a call to population managers and researchers to contact the authors with details of datasets that they feel might contribute information to a general analysis to test the bigger is better paradigm for *S. salar* smolts in the manner described. This is important because a better understanding of how intrinsic and extrinsic factors affect the vital rates of the individuals that constitute a population could allow these populations to be better managed. In the case of *S. salar*, for example, evidence of a general positive effect of smolt size on their subsequent marine survival could support management strategies that maximise not just the number but also the body size and condition of emigrating smolts, perhaps by improving overwintering habitat. (We acknowledge that a management strategy designed to maximise both number and size and condition of smolts would have to account for many complicating factors, such as any negative effect of density dependence on body size.) This could be a particularly pertinent message at present given evidence that body sizes of juvenile *S. salar* are decreasing in countries such as England and France (e.g., Gregory *et al.*, 2017). In summary, determining the role of smolt body size in marine survival could provide considerable conservation and fishery benefits for *S. salar* and could be incorporated into methods currently used to set conservation limits and fishing quotas (MacLean *et al.*, 2003).

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#### References

Amiro, P. G. 2003. Trends in marine survival of hatchery-origin Atlantic salmon (*Salmo salar* L.) to four maritime rivers with comparisons to indices of wild-origin smolt survival to the LaHave river, Nova Scotia, Canada. In *Marine mortality of Atlantic salmon, Salmo salar L: methods and measures* (Potter, E. C. E.; Maoileidigh, N. O. & Chaput, G., eds), pp. 44-58. Canadian Science Advisory Secretariat Research Document 2003/101.

Aprahamian, M. W., Martin Smith, K., McGinnity, P., McKelvey, S. & Taylor, J. (2003). Restocking of salmonids – opportunities and limitations. *Fisheries Research* **62**, 211-227. https://dx.doi.org/10.1016/S0165-7836(02)00163-7.

Armstrong, J. D., Mckelvy, S., Smith, G. W., Rycroft, P. & Fryer, R. J. (*this issue*). Effects of individual variation in length, condition and run-time on return rates of wild-reared Atlantic salmon smolts. *Journal of Fish Biology*.

Auger-Méthé, M, Field, C., Albertsen, C. M., Derocher, A. E., Lewis, M. A., Jonsen, I. D. & Flemming, J. M. (2016). State-space models' dirty little secrets: even simple linear Gaussian models can have estimation problems. *Scientific Reports* **6**, 26677. https://dx.doi.org/10.1038/srep26677. Beaugrand, G & Reid, P. C. (2012). Relationships between North Atlantic salmon, plankton, and hydroclimatic change in the Northeast Atlantic. *ICES Journal of Marine Science* **69**, 1549-1562. https://dx.doi.org/10.1093/icesjms/fss153.

Bentley, N. (2015). Data and time poverty in fisheries estimation: potential approaches and solutions. ICES Journal of Marine Science **72**, 186-193. https://dx.doi.org/10.1093/icesjms/fsu023.

Brook, B. W., Sodhi, N. S. & Bradshaw, C. J. A. (2008). Synergies among extinction drivers under global change. *Trends in Ecology & Evolution* **23**, 453-460. https://dx.doi.org/10.1016/j.tree.2008.03.011.

Caron, F. & Dodson, J. (2003). Change in size selective mortality in Atlantic salmon (*Salmo salar L.*), de la Trinité river, Québec, Canada. In *Marine mortality of Atlantic salmon, Salmo salar L: methods and measures* (Potter, E. C. E.; Maoileidigh, N. O. & Chaput, G., eds), pp. 118-123. Canadian Science Advisory Secretariat Research Document 2003/101.

Clutton-Brock, T & Sheldon, B. C. (2010). Individuals and populations: the role of long-term, individual-based studies of animals in ecology and evolutionary biology. *Trends in Ecology & Evolution* **25**, 562-573. https://dx.doi.org/10.1016/j.tree.2010.08.002.

Dempson, J.; Mullins, C.; Bourgeois, C.; O'Connell, M. & Reddin, D. (2003) Perspectives on smolt production and marine survival of Newfoundland Atlantic salmon (*Salmo salar* L.) related to smolt size and run timing. *In* Potter, E. C. E.; Maoileidigh, N. O. & Chaput, G. (Eds.) Marine mortality of Atlantic salmon, *Salmo salar* L: methods and measures

Dieperink, C., Pedersen, S. & Pedersen, M. I. (2001). Estuarine predation on radiotagged wild and domesticated sea trout (*Salmo trutta* L.) smolts. *Ecology of Freshwater Fish* **10**, 177-183. https://dx.doi.org/10.1034/j.1600-0633.2001.100307.x.

Dieperink, C., Bak, B. D., Pedersen, L.-F., Pedersen, M. I. & Pedersen, S. (2002). Predation on Atlantic salmon and sea trout during their first days as postsmolts. *Journal of Fish Biology* **61**, 848-852. https://dx.doi.org/10.1111/j.1095-8649.2002.tb00917.x.

Francis, R. I. C. C. (1990). Back-calculation of fish length: a critical review. *Journal of Fish Biology* **36**, 883-902. https://dx.doi.org/10.1111/j.1095-8649.1990.tb05636.x.

Gimenez, O., Rossi, V., Choquet, R., Dehais, C., Doris, B., Varella, H., Vila, J.-P. & Pradel, R. (2007). State-space modelling of data on marked individuals. *Ecological Modelling* **206**, 431-438. https://dx.doi.org/10.1016/j.ecolmodel.2007.03.040.

Gregory, S. D., Nevoux, M., Riley, W. D., Beaumont, W. R. C., Jeannot, N., Lauridsen, R. B., Marchand, F., Scott, L. J. & Roussel, J.-M. (2017). Patterns on a parr: drivers of long-term salmon parr length in U.K. and French rivers depend on geographical scale. *Freshwater Biology* **62**, 1117-1129. https://dx.doi.org/10.1111/fwb.12929.

Guillera-Arroita, G. (2017). Modelling of species distributions, range dynamics and communities under imperfect detection: advances, challenges and opportunities. *Ecography* **40**, 281-295. https://dx.doi.org/10.1111/ecog.02445.

Henderson, M. A. & Cass, A. J. (1991). Effect of smolt size on smolt-to-adult survival for Chilko Lake sockeye salmon (*Oncorhynchus nerka*). *Canadian Journal of Fisheries and Aquatic Sciences* **48**, 988-994. https://dx.doi.org/10.1139/f91-115.

Holtby, L. B., Andersen, B. C. & Kadowaki, R. K. (1990). Importance of smolt size and early ocean growth to interannual variability in marine survival of Coho salmon (*Oncorhynchus kisutch*). *Canadian Journal of Fisheries and Aquatic Sciences* **47**, 2181-2194. https://dx.doi.org/10.1139/f90-243.

Jonsson, B., Jonsson, M. & Jonsson, N. (2017). Influences of migration phenology on survival are size dependent in juvenile Atlantic salmon (*Salmo salar*). *Canadian Journal of Zoology* (online) https://dx.doi.org/10.1139/cjz-2016-0136.

Jutila, E., Jokikokko, E. & Julkunen, M. (2006). Long-term changes in the smolt size and age of Atlantic salmon, *Salmo salar* L., in a northern Baltic river related to parr density, growth opportunity and postsmolt survival. *Ecology of Freshwater Fish* **15**, 321-330. https://dx.doi.org/10.1111/j.1600-0633.2006.00171.x.

Kallio-Nyberg, I., Jutila, E., Saloniemi, I. & Jokikokko, E. (2004). Association between environmental factors, smolt size and the survival of wild and reared Atlantic salmon from the Simojoki River in the Baltic Sea. *Journal of Fish Biology* **65**, 122-134. https://dx.doi.org/10.1111/j.0022-1112.2004.00435.x.

Kéry, M. & Schaub, M. (2011). *Bayesian population analysis using WinBUGS: a hierarchical perspective*. New York, NY: Academic Press.

Koenings, J. P., Geiger, H. J. & Hasbrouck, J. J. (1993). Smolt-to-adult survival patterns of Sockeye salmon (*Oncorhynchus nerka*): effects of smolt length and geographic latitude when entering the sea. *Canadian Journal of Fisheries and Aquatic Sciences* **50**, 600-611. https://dx.doi.org/10.1139/f93-069.

Limburg, K. E. & Waldman, J. R. (2009). Dramatic declines in North Atlantic diadromous fishes. *BioScience* **59**, 955-965. https://dx.doi.org/10.1525/bio.2009.59.11.7.

MacLean, J.; Hansen, L. & Friedland, K. (2003) Links between environment, growth and survival in Atlantic salmon (*Salmo salar* L.) and the implications for modelling prefishery abundance. In *Marine mortality of Atlantic salmon, Salmo salar L: methods and measures* (Potter, E. C. E.; Maoileidigh, N. O. & Chaput, G., eds), pp. 137-149. Canadian Science Advisory Secretariat Research Document 2003/101.

Mills, K. E., Pershing, A. J., Sheehan, T. F. & Mountain, D. (2013). Climate and ecosystem linkages explain widespread declines in North American Atlantic salmon populations. *Global Change Biology* **19**, 3046-3061. https://dx.doi.org/10.1111/gcb.12298.

Molony, B. W., Lenanton, R., Jackson, G. & Norriss, J. (2003). Stock enhancement as a fisheries management tool. *Reviews in Fish Biology and Fisheries* **13**, 409-432. https://dx.doi.org/10.1007/s11160-004-1886-z.

Newton, M., Barry, J., Dodd, J. A., Lucas, M. C., Boylan, P. & Adams, C. E. (2016). Does size matter? A test of size-specific mortality in Atlantic salmon *Salmo salar* smolts tagged with acoustic transmitters. *Journal of Fish Biology* **89**, 1641-1650. https://dx.doi.org/10.1111/jfb.13066.

Otero, J., L'Ábee-Lund, J. H., Castro-Santos, T., Leonardsson, K., Storvik, G. O., Jonsson, B., Dempson, D., *et al.* (2014). Basin-scale phenology and effects of climate variability on global timing of initial seaward migration of Atlantic salmon (*Salmo salar*). *Global Change Biology* **20**, 61-75. https://dx.doi.org/10.1111/gcb.12363.

Parent, E. & Rivot, E. (2012). *Introduction to hierarchical Bayesian modeling for ecological data*. New York, NY: CRC Press.

Ricker, W. E. (1976). Review of the rate of growth and mortality of Pacific salmon in salt water, and noncatch mortality caused by fishing. *Journal of the Fisheries Research Board of Canada* **33**, 1483-1524. https://dx.doi.org/10.1139/f76-191.

Riley, W. D., Ibbotson, A. T., Beaumont, W. R. C., Pawson, M. G., Cook, A. C. & Davison, P. I. (2011). Predation of the juvenile stages of diadromous fish by sea bass (*Dicentrarchus labrax*) in the tidal reaches of an English chalk stream. *Aquatic Conservation: Marine and Freshwater Ecosystems* **21**, 307-312. https://dx.doi.org/10.1002/aqc.1184.

Robinson, R. A., Morrison, C. A. & Baillie, S. R. (2014). Integrating demographic data: towards a framework for monitoring wildlife populations at large spatial scales. *Methods in Ecology and Evolution* **5**, 1361-1372. https://dx.doi.org/10.1111/2041-210X.12204.

Royle, J. A. & Dorazio, R. M. (2008). *Hierarchical modeling and inference in ecology: the analysis of data from populations, metapopulations and communities*. New York, NY: Academic Press.

Russell, I. C., Aprahamian, M. W., Barry, J., Davidson, I. C., Fiske, P., Ibbotson, A. T., Kennedy, R. J., Maclean, J. C., Moore, A. & Otero, J. (2012). The influence of the freshwater environment and the biological characteristics of Atlantic salmon smolts on their subsequent marine survival. *ICES Journal of Marine Science* **69**, 1563-1573. https://dx.doi.org/10.1093/icesjms/fsr208.

Salminen, M., Kuikka, S. & Erkamo, E. (1995). Annual variability in survival of sea-ranched Baltic salmon, *Salmo salar* L: significance of smolt size and marine conditions. *Fisheries Management and Ecology* **2**, 171-184. https://dx.doi.org/10.1111/j.1365-2400.1995.tb00110.x.

Saloniemi, I., Jokikokko, E., Kallio-Nyberg, I., Jutila, E. & Pasanen, P. (2004). Survival of reared and wild Atlantic salmon smolts: size matters more in bad years. *ICES Journal of Marine Science* **61**, 782-787. https://dx.doi.org/10.1016/j.icesjms.2004.03.032.

Sogard, S. M. (1997). Size-selective mortality in the juvenile stage of teleost fishes: a review. *Bulletin of Marine Science* **60**, 1129-1157.

## **Electronic References**

ICES. (2017). Report of the Working Group on North Atlantic Salmon (WGNAS) 29 March–7 April 2017, Copenhagen, Denmark. Available at http://ices.dk/sites/pub/Publication%20Reports/Expert%20Group%20Report/acom/2017/WG

NAS/wgnas\_2017.pdf (last accessed 6 August 2017).

# Tables

Table 1: Studies (non-exhaustive) testing the effect of salmonid smolt length on the subsequent marine survival, with a focus on Salmo salar. Origin: W =

wild, $H =$ hatchery.	Hw = hatchery	from wild s	stock; Number	of smolts: NI	R = not reported.
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Туре	Method	Study	Species	Origin	Evidence	Number	Number	Number	Unit of	Data characteristics	Model characteristics
						smolts	years	rivers	variance		
1	Relating annual mean length	Amiro (2003)	Salmo salar	W Hw	No effect of length on adult	NR	4	1	Year	Small sample size,	Linear only, No covariates
	to annual overall adult return				return rate					Imperfect adult detection	
	rate										
1	Relating annual mean length	Armstrong et	Salmo salar	Hw	Positive effect of length on	>12500	15	1	Individual	Medium sample size,	Linear only, Individual-
	to 1SW and 2+SW adult	al. (2017)			1SW and 2+SW adult return					Long time series, Perfect	level covariates
	return rates				rate					adult detection	
2	Relating annual lengths back-	Caron &	Salmo salar	W	Positive effect of length on	> 262000	4	1	Individual	Large sample size, Short	No covariates, Unaccounted
	calculated from 1SW and	Dodson (2003)			1SW and 2SW adult return					time series	back-calculation uncertainty
	2SW returned adult scales to				rate						
	annual lengths										
1	Relating mean length and	Dempson, et al.	Salmo salar	W	Weak positive effect of length	>1300000	10	6	River	Multiple rivers for	Linear only, No covariates
	adult return rate	(2003)			on adult return rate					generality, Small sample	
										size, Perfect adult	
										detection	
1	Relating annual mean length	Dempson, et al.	Salmo salar	W	Weak positive and negative	12849-404667	9-16	1	Year	Medium sample size,	Linear only, No covariates
	and annual adult return rate	(2003)			river-specific effects of length					Long time series, Perfect	
					on adult return rate					adult detection	
3	Retrospective classification	Dieperink, et	Salmo trutta	WΗ	Weak positive effect of length	37	1	1	Binomial	Small sample size,	No covariates, Unaccounted
	of radio-tagged individual	al. (2001)			on probability to be predated				error	Imperfect smolt	classification uncertainty,

	fates during their early				by bird					detection	Potential confounding by
	(estuarine) seaward migration										origin
3	Retrospective classification	Dieperink, et	Salmo salar &	W	Weak positive effect of length	24 (S. salar) &	1	1	Binomial	Small sample size,	No covariates, Unaccounted
	of radio-tagged individual	al. (2002)	Salmo trutta		on probability to be predated	15 (S. trutta)			error	Imperfect smolt	classification uncertainty
	fates during their early				by bird					detection	
	(estuarine) seaward migration										
1	Relating annual mean length	Henderson &	Oncsrhynchus	W	No effect of length on adult	NR	34	1	Year	Medium sample size,	Linear only, No covariates
	to annual overall adult return	Cass (1991)	nerka		return rate					Long time series,	
	rate									Imperfect adult detection	
2	Relating annual mean length	Henderson &	Oncsrhynchus	W	Significantly higher mean	585, 474, 484	3	1	Individual	Imperfect detection	Separate years,
	back-calculated from returned	Cass (1991)	nerka		length back-calculated from						Unaccounted back-
	adult scales to annual mean				returned adult scales for 2						calculation uncertainty, No
	lengths				years						covariates
2	Relating annual lengths back-	Holtby, et al.	Oncsrhynchus	W	Significantly higher(lower)	NR	14	1	Individual	Imperfect detection	Separate years,
	calculated from returned adult	(1990)	kisutch		mean length back-calculated						Unaccounted back-
	scales to annual lengths				from returned adult scales for						calculation uncertainty, No
					7(2) years; equal in 5 years						covariates
4	Comparing mean marine	Jonsson, et al.	Salmo salar	W	Higher survival among longer	36833	37	1	Group	Large sample size, Long	Unaccounted tag mortality,
	survival among length classes	(2017)			individuals migrating during					time series, Perfect adult	No formal statistical test,
	migrating in different time				middle emigration period					detection	Group-level covariates
	periods										
1	Relating annual mean length	Jutila, et al.	Salmo salar	Hw	Positive effect of length on	NR	23	1	Year	Large sample size, Long	Linear only, Unaccounted
	to annual tag recovery rate	(2006)			post-smolt tag recovery rate					time series	tag mortality, Unknown
											reporting effort, No
											covariates
1	Relating annual mean length	Kallio-Nyberg,	Salmo salar	Hw	No evidence of size-dependent	>15000	21	1	Year	Large sample size, Long	Linear only, Unaccounted
	to annual tag recovery rate	et al (2004)			mortality					time series	tag mortality, Unknown

											reporting effort, No
											covariates
1	Relating annual mean length	Koenings, et al.	Oncorhynchus	WΗ	Positive effect of length on	NR	1-9	12	Year	Multiple rivers for	Linear and non-linear,
	to annual overall adult return	(1993)	nerka		adult return rate, with possible					generality, Medium	Unaccounted river effect,
	rate				quadratic effect					sample size, Imperfect	River-level covariates
										adult detection	
3	Acoustic-tagged individual	Newton, et al.	Salmo salar	W	No evidence of size-dependent	68	2	1	Individual	Imperfect detection	Separate years, No
	lengths compared to their	(2016)			mortality						covariates
	fates during early (estuarine)										
	seaward migration										
4	Early (estuarine) marine tag	Salminen, et al.	Salmo salar	Н	Positive and no effects of	35000-505000	12	2	Year	Large sample size, Long	Separate rivers,
	recovery rate calculated and	(1995)			length on tag recovery rate	& 11000-				time series, Imperfect	Unaccounted tag mortality,
	plotted for different smolt					577000				detection	Unknown reporting effort,
	size classes										No formal statistical test
1	Relating individual length to	Saloniemi, et	Salmo salar	WΗ	Positive effect of length on tag	>3740	2	1	Individual	Medium sample size,	Linear only, Unaccounted
	early (estuarine) marine tag	al. (2004)			recovery rate					Short time series,	tag mortality, Unknown
	recovery rate									Imperfect detection	reporting effort, Group-
											level covariates

### **Figure Captions**

Figure 1: Atlantic salmon populations are declining, as suggested by the 5-year rolling mean reported nominal catch (tonnes). Source: ICES2017.

Figure 2: Fitted effect of *S. salar* smolt length on their subsequent marine survival. (a) effect measured using a cumulative link mixed model including covariates condition and day and a random year effect. The dots show the observed proportions of returning fish; these are calculated by splitting the marginal distribution of the explanatory variable (length) into twenty bins with equal numbers of fish and calculating the proportion of returning fish in each. The position of the dots on the x-axis are the 2.5, 7.5, ..., 97.5 percentiles of the explanatory variable so, 95% of fish had lengths between 104 and 143 mm. The solid lines span the observed range of each explanatory variable. The full range of lengths illustrated by the fitted line. Models with smooth relationships were also explored, but the nonlinear terms were never significant. Redrawn from Armstrong *et al.* (2017). (b) effect measured using a logistic regression including covariates origin and year and using just two years of data: black lines = 1991, grey lines = 1993, solid lines = wild smolts, dashed lines = reared smolts. Redrawn from Saloniemi *et al.* (2004). Note the similarity in intercept and slopes of the fits.

## Figures



Fig. 1









Fig. 2