

Program SampleSize 3

Sample Size Calculations for Fish and Wildlife Survival Studies

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Chapter 1: Introduction

Release-recapture studies are commonly used to estimate survival-related parameters in fish and wildlife studies. In designing such a study, one must determine the quantity of tagged individuals needed to achieve the desired precision without wasting resources by using larger release sizes than necessary. Program SampleSize was developed to fill this need. It allows the user to input a range of values for a release size or estimated parameters and calculate the resulting precision of survival projections. It also graphically displays the calculated precision as a function of the range of values, allowing the user to observe the sensitivity of the precision over the range.

There are seven study design types implemented in Program SampleSize:

- 1. Single Site,
- 2. Single Release,
- 3. Paired Release,
- 4. Virtual Paired Release,
- 5. Ricker Two-Release,
- 6. ViRDCt (Full model), and
- 7. ViRDCt (Reduced model).

Note that previous versions of Program SampleSize included two study types not included in the current version: (1) transport in-river ratio and (2) balloon-tag passage survival estimate. These study types are no longer explicitly included because the transport in-river ratio study is an example of a Ricker study, and the variance for a balloon-tag study is the same as the variance for the Ricker model.

Chapter 2 describes the basics in using Program SampleSize, version 3.0. The subsequent chapters describe the details in using Program SampleSize for the specific models mentioned above.

Chapter 2: Using Program SampleSize

Figure 1 shows Program SampleSize at startup, displaying the "Welcome" screen.



Figure 1. Program SampleSize at startup, displaying the Welcome screen.

On the left side under the heading "Study Design Selection," the user can click on any one of the seven study design types. Clicking on one of the selections will bring up a tabbed dialog for that study design.

Figure 2 shows the program after clicking on "Single Release" under "Study Design Types," bringing up the tabbed window for a Single Release release-recapture study design.

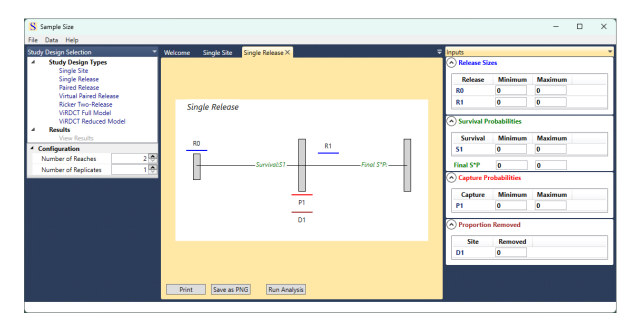


Figure 2. Program SampleSize with the Single Release study design selected.

The user may have more than one study design tab open at a time. In Figure 3, the user has both a Single Release study design and a Paired Release study design open, with the Paired Release tab currently active.



Figure 3. Program SampleSize with multiple study designs open.

The user may close the current tab by clicking the "X" on the tab. They may also open the context menu for other options by right-clicking on the tab as shown in Figure 4.

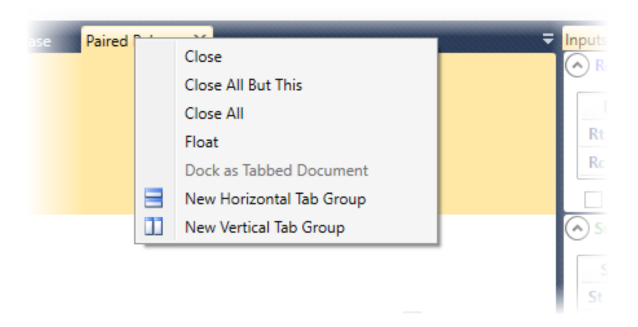


Figure 4. The context menu for the study design tabs.

On each study tab, there are three buttons at the bottom (shown in Figure 5) that allow the user to print the study diagram, save it as a .png file, or run the analysis once all inputs have been entered.

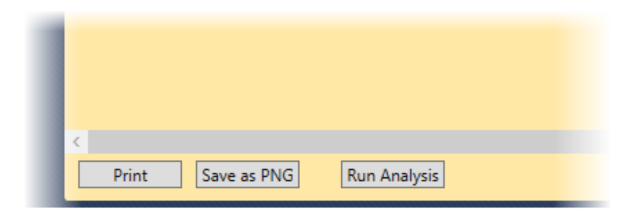


Figure 5. The buttons at the bottom of a study design diagram for printing or saving the diagram and for running the analysis.

Once a study tab is active, one or two additional areas appear on the SampleSize dialog: (1) a "Configuration" section, if applicable, under the "Study Design Selection" section on the left and (2) the "Inputs" section on the right.

Figure 6 shows the Configuration section for a Single Release study design. In this case, the user may specify the number of reaches and the number of replicates (explained in Chapter 4).

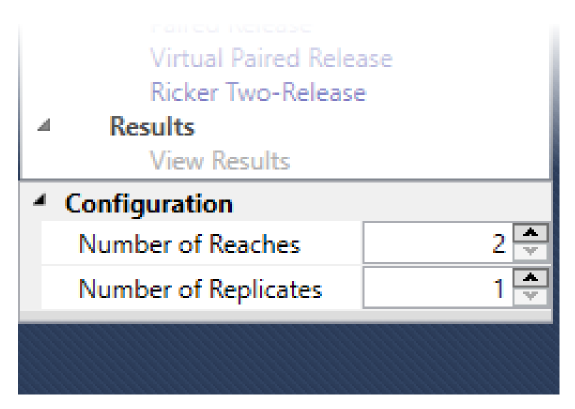


Figure 6. The Configuration menu for a Single Release study design.

2.1 Input Section

Figure 7 shows the "Inputs" section for a Single Release study design. In this section, the user inputs all release sizes, estimates of survival probabilities, capture probabilities, and the proportion removed from each site (when applicable). Release sizes must be integers, and survival and capture probabilities as well as proportions removed must be values between 0.0 and 1.0.

Most inputs have a place to enter both a minimum and a maximum value — R0, R1, S1, and P1 in Figure 7. This allows the user to explore the sensitivity of the resulting precision to the input value with a range. Note that only one input can have a range; when both a minimum and maximum for a given input are entered, the "Maximum" input box disappears for the other inputs.

In order to cancel a range for an input, the user must enter a "0" (zero) in the Maximum entry box for the input.

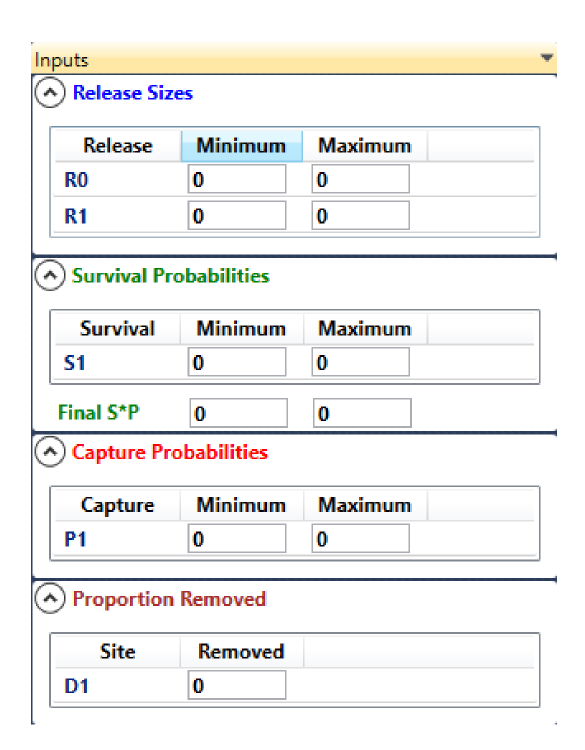


Figure 7. The Inputs section for a Single Release study design.

2.2 File Operations

The "File" menu on the upper left of Program SampleSize allows the user to save the current configuration for all study designs that are open and to restore them from a file at a later time.

2.3 Analysis Results

2.3.1 Analysis with No Range

Figure 8 shows the study design diagram and input section filled in for a Single Release study design without a range specified. Note that the input values are displayed on the diagram upon being entered by the user.

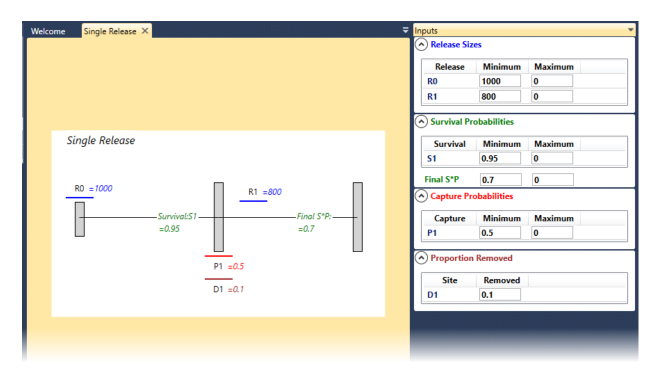


Figure 8. The Single Release study design diagram and Inputs section with no range specified.

When the "Run Analysis" button is pressed, a report appears in a separate window as shown in Figure 9 that lists all user-supplied inputs and a results table with:

- 1. the standard error of the parameter of interest (S1 in this case),
- 2. the half-width of a 95% confidence interval, and
- 3. the half-width of a 90% confidence interval.

At the bottom of the report are two buttons that allow the user to print the report or save it as a comma-separated value (.csv) file.

2.3.2 Analysis with a Range

Figure 10 shows the inputs with a range entered for *R*0; when the range is entered, the Maximum entry box for the other inputs disappear.

When the Run Analysis button is pressed, a results window appears as shown in Figure 11. There are two tabs on the results window when a range is entered: (1) the "Results Table" and (2) the "Results Graph."

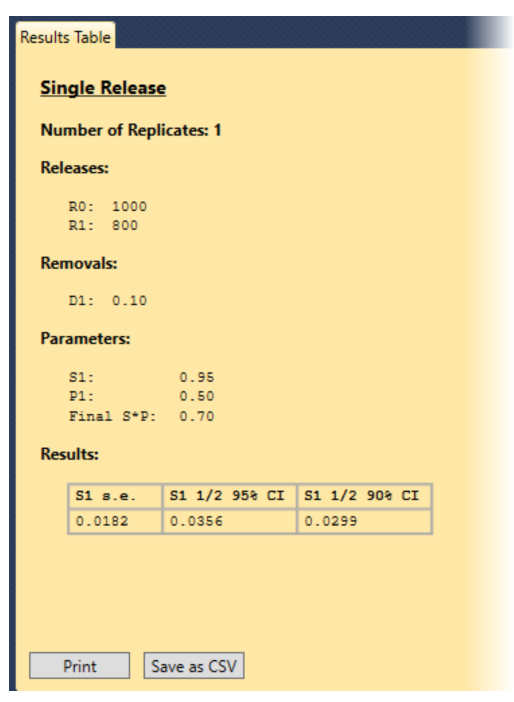


Figure 9. The Results Table window when no range is specified.

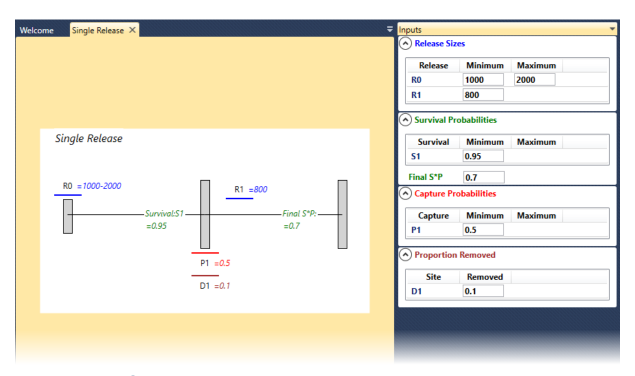


Figure 10. The Single Release study diagram and Inputs section when a range is specified.

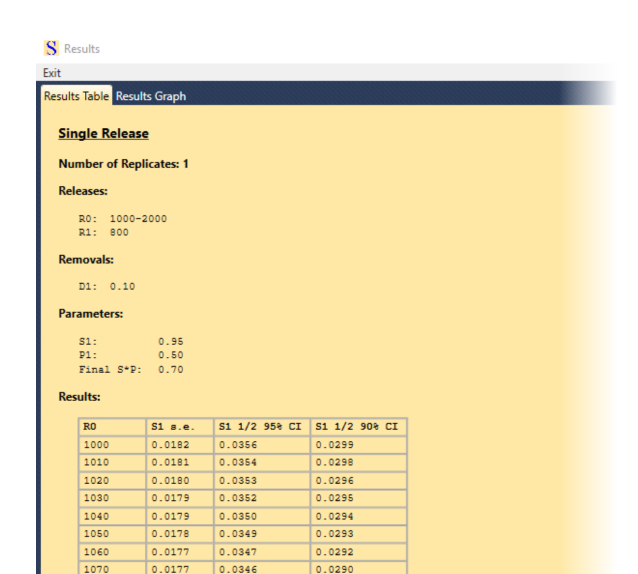


Figure 11. The Results Table window when a range is specified.

0.0289

0.0288

0.0287

0.0286

2.3.3 Results Table

0.0176

0.0175

0.0174

0.0174

0.0345

0.0343

0.0342

0.0341

1080

1090

1100

The "Results Table" tab is active by default when the results window appears. It is similar to the results report when no range is entered, except there is an additional column in the table for the values of the input with a range. The Results Table shows the results for each value of the input along one hundred evenly spaced values along the range. In the table in Figure 11, a range of

1,000–2,000 was entered for *R*0, thus the results table shows results for *R*0 in values that increment by ten.

2.3.4 Results Graph

Figure 12 shows the "Results Graph" tab. The input with the range (*R*0 in this case) is on the *X*-axis and the half-width of the confidence interval (CI) is on the Y-axis. The red line (on top in Figure 12) represents a 95% CI and the blue line represents a 90% CI. When the user moves the mouse along the graph horizontally, a vertical line follows the mouse and shows CI half-width values for 95% and 90%. The user can click on any point on the graph and a fixed line will appear showing the CI half-width values at that point.

In Figure 12, the user has clicked on the value R0 = 1,400, and a line has appeared showing the half-width of a 95% CI as 0.0309 and the half-width of a 90% CI as 0.0259. The mouse currently rests over the value R0 = 1710.

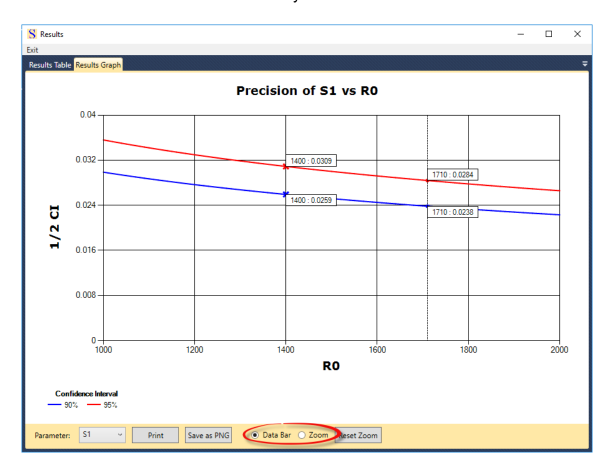


Figure 12. The Results Graph.

At the bottom of the graph are two option buttons (circled in Figure 12) labeled "Data Bar" and "Zoom." The Data Bar button is selected by default, enabling the behavior just described. If the user clicks on Zoom, the user can hold the left mouse button and drag over a portion of the graph in order to focus on that section. In Figure 13, the user has dragged the mouse over a portion of the graph indicated in gray between R0 = 1200 and R0 = 1600. When the mouse button is released, the Results Graph shows only that portion, as shown in Figure 14. The user can restore the graph to its initial state by clicking on the "Reset Zoom" button at the bottom.

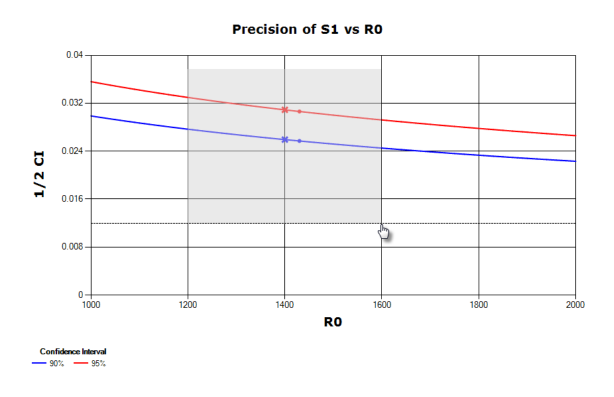


Figure 13. The user selecting a section of the Results Graph to zoom in on.

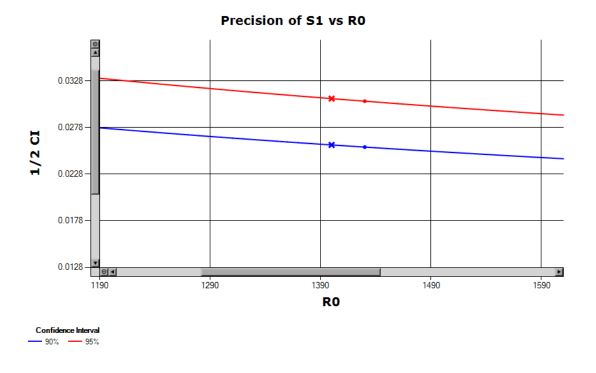


Figure 14. The Results Graph after the user the user has zoomed in on a section.

At the bottom are buttons to print the graph or to save it as a .png file. There is also a box labeled "Parameter" showing the parameter of interest that the CIs apply to. For a Single Release design only, this is a combo box where the user can select the survival parameter of interest (described in Chapter 4).

Chapter 3: Single Site Study Design

The Single Site study design is the simplest of the available designs, using only one release and one downstream detection site. Figure 15 shows the layout of the Single Site study design in Program SampleSize. There is only one estimable parameter: the joint probability of survival to, and detection at, the downstream site ("Final $S \times P$ "). The two inputs are the release size (R) and the Final $S \times P$ parameter.

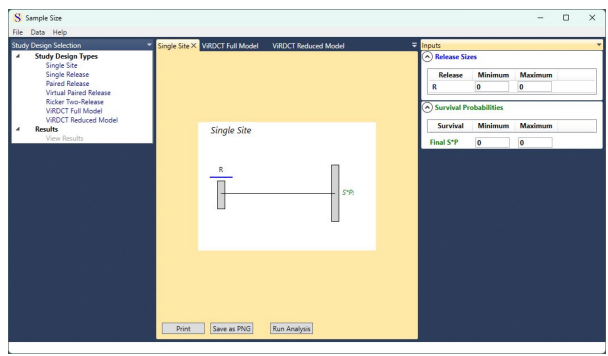


Figure 15. The Single Site study diagram.

Chapter 4: Single Release Study Design

The Single Release study design differs from the Single Site study design by using more than one detection site, providing the data necessary to estimate survival in all reaches except the final reach. Multiple replicates are allowed, as well. When the Single Release study design is selected, the user can specify the number of reaches and the number of replicates under the "Configuration" heading at the left of the SampleSize dialog.

In Figure 16, the user has selected 4 reaches and 2 replicates. The diagram and inputs are adjusted for the number of reaches; the resulting variance calculations will be adjusted for the number of replicates specified.

When a diagram becomes too large to show in entirety (e.g., when adding more reaches in the Single Release design), a scroll bar appears at the bottom, allowing the user to see the rest of the diagram.

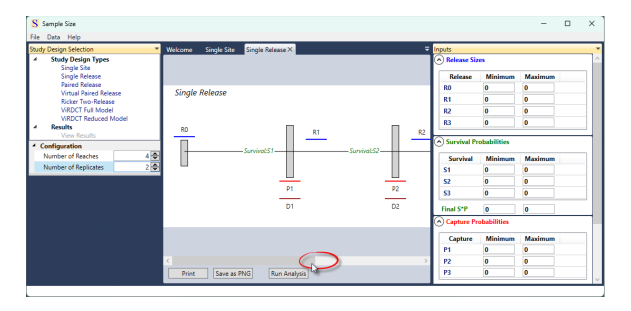
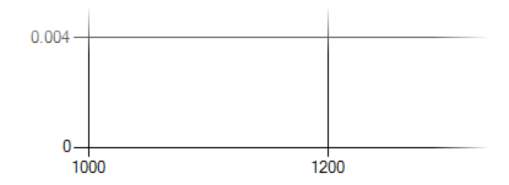


Figure 16. The Single Release study diagram with a scroll bar at the bottom.

4.1 Single Release Results Graph

When there are three or more reaches in a Single Release design, the user can select the survival parameter of interest. Figure 17 shows the lower left portion of the Results Graph for a Single Release design with four downstream reaches and three survival parameters: *S*1, *S*2, and *S*3. The user can select the parameter of interest, and the graph will display the CI half-width values for the chosen parameter.



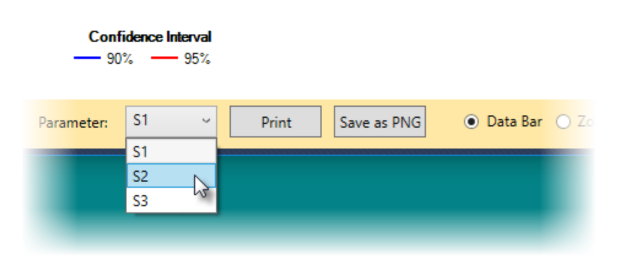


Figure 17. Selecting the parameter of interest on the Results Graph for a Single Release study design with four reaches.

Chapter 5: Paired Release Study Design

Figure 18 shows the diagram for a Paired Release study design. As with the Single Release study design, the user can specify the number of reaches and the number of replicates. Unlike the Single Release study design, the results will only show the CI half-width values for the treatment survival (*St*).

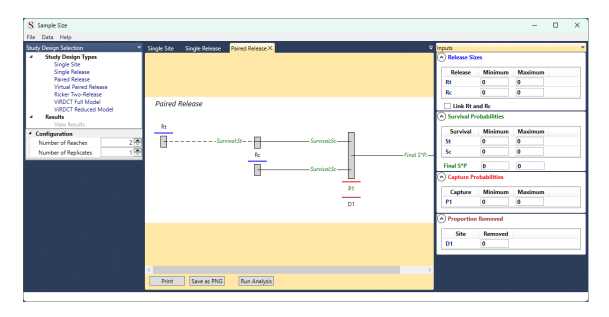


Figure 18. The Paired Release study design diagram.

Program SampleSize allows the user to equate the sizes of the treatment release and control release under the Inputs section as shown in Figure 19. When the "Link *Rt* and *Rc*" check box is checked, the control release size value (*Rc*) will be linked (equated) to the value of the treatment release (*Rt*). If a range is then specified for *Rt*, then both the treatment and control release sizes will be varied together over the range.

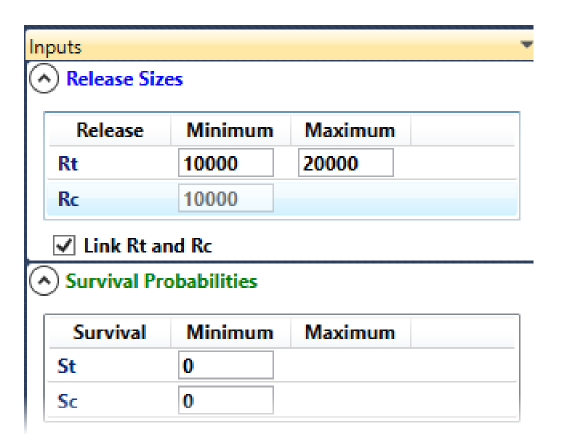


Figure 19. Linking the treatment and control releases in a Paired Release study design.

Chapter 6: Virtual Paired Release Study Design

Figure 20 shows the diagram for a Virtual Paired Release study design. There are no configuration options for a Virtual Paired Release study design, and the parameter of interest for which CIs will be calculated is the dam passage survival (*Sdam*). Analogous to the Paired Release study design, releases *R*2 and *R*3 can be linked using the same action as linking *Rt* and *Rc* in the Paired Release study design (Figure 21).

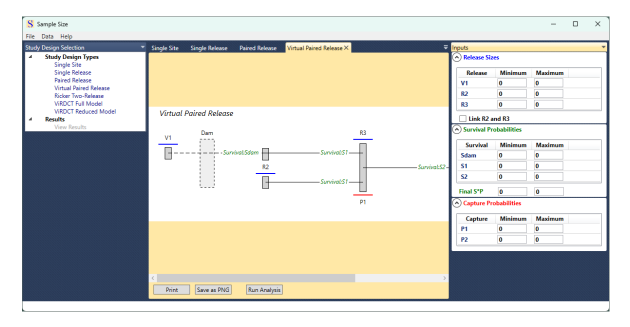


Figure 20. The Virtual Paired Release study diagram.

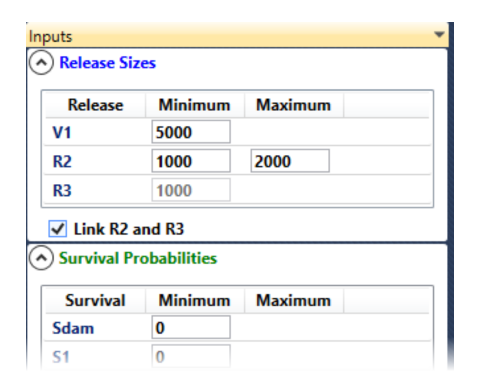


Figure 21. Linking releases *R*2 and *R*3 in a Virtual Paired Release study design.

Chapter 7: Ricker Two-Release Study Design

Figure 22 shows the diagram for the Ricker Two-Release study design. There are no configuration options, and the parameter of interest for which CIs are calculated is the treatment survival (*St*). The treatment and control releases (*Rt* and *Rc*) share the same downstream capture probability, *P*, and can be further linked by equating their release sizes.

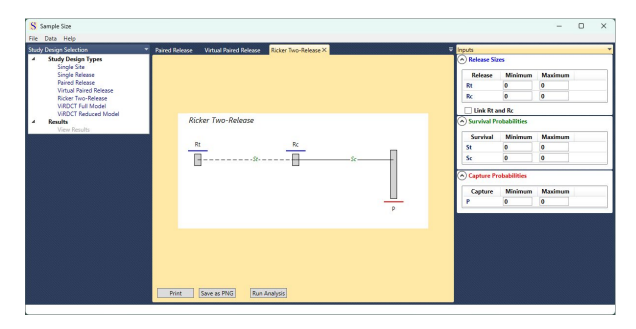


Figure 22. The Ricker Two-Release study diagram.

The survival probability from release to the detection site is $St \times Sc$ for the treatment release (Rt) and is Sc for the control release (Rc). Thus, the treatment survival (St) is calculated as the ratio of the two survival probabilities to the detection site.

Chapter 8: ViRDCt Study Design (Full Model)

Figure 23 shows the diagram for the ViRDCt Full Model study design. There are no configuration options, and the parameter of interest for which CIs are calculated is the dam passage survival (*SD*).

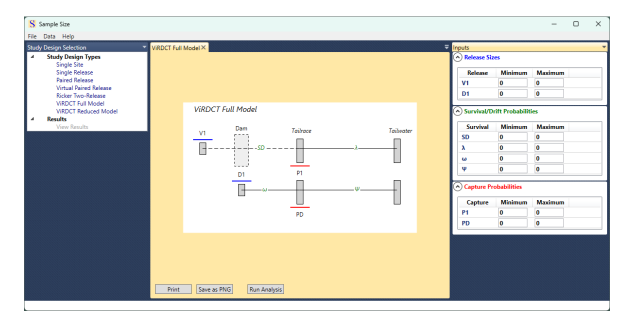


Figure 23: The "ViRDCt Full Model" study diagram.

There are two releases for the ViRDCt model:

Parameter Definition

V1 = virtual release, consisting of fish detected at the dam face,and

D1 = dead fish released in the dam tailrace.

The parameters for the full ViRDCt model are as follows:

Parameter Definition

- SD = dam passage survival,
 - λ = join probability of survival between tailrace and tailwater arrays and being detected at the tailwater array,
 - ω = probability of a dead fish from *D*1 arriving at the tailrace array,
- Ψ = joint probability that a dead fish is washed down to the tailwater array from the tailrace array and is detected at the tailwater array,
- P1 = probability of an alive V1 fish being detected at the tailrace array, and
- *PD* = probability of detecting a dead fish at the tailrace array.

Chapter 9: ViRDCt Study Design (Reduced Model)

Figure 24 shows the diagram for the ViRDCt Reduced Model study design. There are no configuration options, and the parameter of interest for which CIs are calculated is the dam passage survival (*SD*).

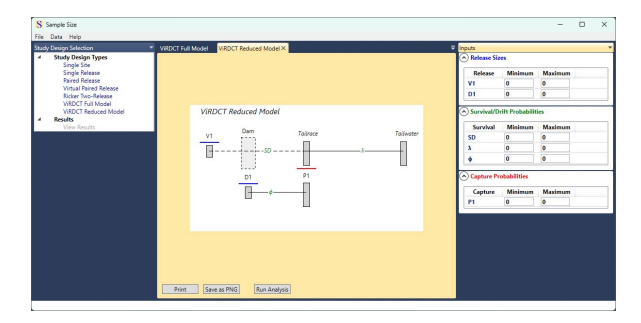


Figure 24: The ViRDCt Reduced Model study diagram.

The reduced model study design has two releases, *V*1 and *D*1 as described in the previous section for the full ViRDCt model study design. The parameters for the reduced ViRDCt model are:

Parameter Definition

- SD = dam passage survival,
 - λ = joint probability of survival between tailrace and tailwater arrays, and being detected at the tailwater array,
 - φ = joint probability of a dead released fish (D1) arriving at the
 tailrace array and being detected at that array, and
- P1 = probability of an alive V1 fish being detected at the tailrace.

Appendix: Variance Calculations

A.1: Single Site Study Design

The only parameter has a binomial distribution, hence:

$$\operatorname{Var}(\widehat{\theta}) = \frac{\theta(1-\theta)}{R},$$

where

Parameter Definition

 $\theta = S \times P$ = joint probability of survival to and detection at the one downstream site, and

R = release size.

A.2: Single Release Study Design

The variance calculations for the Single Release study design are adapted from Burnham et al. (1987).

For survival parameter S_i ,

$$Var(\hat{S}_i) = S_i^2(f_1 + f_2 + f_3)$$
 for $i = 1$ to $K - 1$,

where

Parameter Definition

K = the number of reaches,

$$f_1 = \frac{1}{E(r_{i-1})} - \frac{1}{E(M_{i-1})},$$

$$f_2 = (1 - P_i)^2 \left(\frac{1}{E(r_i)} - \frac{1}{E(M_i)} \right),$$

$$f_3 = P_i(1 - P_i) \frac{(1 - E(A_i))^2}{E(A_i)E(T_i)},$$

and within those equations:

Parameter Definition

$$E(r_i) = E(M_i)(1 - \chi_i),$$

 χ_i = probability of not being detected after occasion i,

$$\chi_{K-1} = 1 - S_K P_K,$$

$$\chi_i = 1 - S_{i+1} + S_{i+1} (1 - P_{i+1}) \chi_{i+1}$$
 for $i = 0$ to $K - 1$,

 M_i = number released from site i, including both those initially released and those previously released, detected and not removed at site i,

$$=R_i+m_i-d_i,$$

 R_i = number initially released from site i,

 m_i = number of previously released individuals detected at site i,

 d_i = number of previously released individuals removed at site i,

$$E(M_i) = R_i +$$

$$\sum_{k=0}^{i-1} R_k \left\{ \prod_{j=k+1}^{i-1} S_j [1 - P_j + P_j (1 - D_j)] \right\} \times S_i P_i (1 - D_i),$$

 P_i = capture probability at site i,

 D_i = proportion removed at site i,

$$E(A_i) = 1 - \chi_i,$$

 T_i = number of previously released individuals detected at or after site i,

$$= m_i + z_i$$
,

 z_i = number of previously released individuals detected after site i but not at site i,

$$=\sum_{j=0}^{i-1}(r_j-m_{j+1}),$$

$$E(T_i) = S_i[P_i + (1 - P_i)(1 - \chi_i)] \times$$

$$\left[\sum_{k=0}^{i-1} R_k \left\{ \prod_{j=k+1}^{i-1} S_j [1 - P_j + P_j (1 - D_j)] \right\} \right],$$

where

$$\prod_{j=k+1}^{k} S_{j} [1 - P_{j} + P_{j} (1 - D_{j})] \equiv 1.$$

For a replicated survival study with n replicates, survival in reach i is estimated by \hat{S}_i :

$$\hat{\bar{S}}_i = \frac{\sum_{j=1}^n \hat{S}_{ij}}{n},$$

where \hat{S}_{ij} is the reach-i survival estimate for replicate j (j=1,...,n). The variance of \hat{S}_i is composed of the population variability in survival for reach i and the average sampling variability for that reach:

$$\operatorname{Var}\left(\hat{\bar{S}}_{i}\right) = \frac{1}{n} \left[\sigma_{S_{i}}^{2} + \frac{\sum_{j=1}^{n} \operatorname{Var}(\hat{S}_{ij} | S_{ij})}{n} \right],$$

where $\sigma_{S_i}^2$ is the natural variation in S_i . The quantity $Var(\hat{S}_{ij}|S_{ij})$ is the measurement error of \hat{S}_{ij} given S_{ij} for replicate j.

In turn,

$$\operatorname{Var}\left(\hat{\bar{S}}_{i}\right) = \frac{\sigma_{S_{i}}^{2}}{n} + \frac{\sum_{j=1}^{n} \operatorname{Var}(\hat{S}_{ij}|S_{ij})}{n^{2}}$$
$$= \frac{\sigma_{S_{i}}^{2}}{n} + \frac{\overline{\operatorname{Var}(\hat{S}_{i}|S_{i})}}{n}.$$

So then

$$SE\left(\hat{\bar{S}}_{i}\right) = \sqrt{\frac{\sigma_{S_{i}}^{2} + \overline{Var(\hat{S}_{i}|S_{i})}}{n}}.$$
 A. 1

If $\sigma_{S_i}^2 = 0$, then

$$SE\left(\hat{\bar{S}}_{i}\right) = \frac{\sqrt{\overline{Var(\hat{S}_{i}|S_{i})}}}{\sqrt{n}} \text{ or } \frac{\sqrt{\sum_{j=1}^{n} Var(\hat{S}_{ij}|S_{ij})}}{n}.$$

If it is further assumed that $Var(\hat{S}_{ij}|S_{ij}) = Var(\hat{S}_{i}|S_{i})$ for all replicates j (j = 1, ..., n), then the expected standard error for reach i becomes

$$SE\left(\hat{S}_{i}\right) = \frac{\sqrt{nVar(\hat{S}_{i}|S_{i})}}{n},$$

$$SE\left(\hat{S}_{i}\right) = \frac{SE(\hat{S}_{i}|S_{i})}{\sqrt{n}}.$$
A. 2

Either Equation (A.1) or (A.2) is correct depending on sample size assumptions.

Note that this development is for sample size calculations rather than for data analysis. For data analysis, a bias correction is necessary because $\widehat{\text{Var}}\left(\widehat{S}_i\right) \neq \text{Var}\left(\widehat{S}_i\right)$.

A.3 Paired Release Study Design

Let $S_1 = S_t S_c$. Then

$$S_t = \frac{S_1}{S_c}.$$

The variances of \hat{S}_1 and \hat{S}_c are calculated as with the Single Release study design in section A.2, treating each release as independent of the other. $Var(\hat{S}_t)$ is then calculated using the delta method and the exact formula for the variance of the product of two independent variables (Seber, 1982):

$$Var(\hat{S}_t) = S_t^2 \left(\frac{Var(\hat{S}_1)}{S_1^2} + \frac{Var(\hat{S}_c)}{S_c^2} + \frac{Var(\hat{S}_1)Var(\hat{S}_2)}{S_1^2 S_c^2} \right).$$

A.4 Virtual Paired Release Study Design

Let

Parameter Definition

$$\phi_1 = Sdam \times S_1$$
,

$$\phi_2 = S_1 S_2$$
, and

$$\phi_3 = S_2$$
.

Then

$$Sdam = \frac{\Phi_1}{\Phi_2/\Phi_3} = \frac{\Phi_1\Phi_3}{\Phi_2}.$$

Treating all three releases as independent from each other, the variances of $\widehat{\varphi}_1$, $\widehat{\varphi}_2$ and $\widehat{\varphi}_3$ are calculated as with the Single Release study design in section A.2. Thus, the variance is calculated using the methods described in section A.3, as follows:

$$Var(\widehat{Sdam}) = \left(\frac{1}{\varphi_2^2} + \frac{Var(\widehat{\varphi}_2)}{\varphi_2^4}\right) \times \left[\varphi_1^2 Var(\widehat{\varphi}_3) + \varphi_3^2 Var(\widehat{\varphi}_1) + Var(\widehat{\varphi}_1) Var(\widehat{\varphi}_3)\right] + \frac{(\varphi_1 \varphi_3)^2}{\varphi_2^4} Var(\widehat{\varphi}_2).$$

A.5 Ricker Two-Release Study Design

$$\operatorname{Var}(\hat{S}_t) = \frac{S_t}{S_c \times P} \left(\frac{(1 - S_c S_t P)}{R_t} + \frac{(1 - S_c P) S_t}{P_c} \right).$$

A.6 ViRDCt Study Design: Full and Reduced Models

For the ViRDCt study design, there are no closed form variance estimates for either the full or reduced model. Thus, the variance is calculated numerically from the likelihood.

The likelihood for the Full ViRDCt model:

$$L = \begin{pmatrix} V_1 \\ n_{11} & n_{01} & n_{10} & n_{00} \end{pmatrix} (S_D \times P_1 \times \lambda + (1 - S_D)\omega P_D \times \psi)^{n_{11}}$$
$$\times (S_D (1 - P_1)\lambda + (1 - S_D)\omega (1 - P_D)\psi)^{n_{01}}$$
$$\times (S_D P_1 (1 - \lambda) + (1 - S_D)\omega P_D (1 - \psi))^{n_{10}}$$

$$\times (S_D(1-P_1)(1-\lambda) + (1-S_D)(1-\omega + \omega(1-P_D)(1-\psi))^{n_{00}}$$

$$\times \begin{pmatrix} D_1 \\ d_{11} & d_{01} & d_{10} \end{pmatrix} \times (\omega P_D \psi)^{d_{11}} \times (\omega(1-P_D)\psi)^{d_{01}}$$

$$\times (\omega P_D(1-\psi))^{d_{10}}(1-\omega + \omega(1-P_D)(1-\psi))^{d_{00}}.$$

The full model has six parameters and six minimum sufficient statistics.

The likelihood for the Reduced ViRDCt model:

$$L = \begin{pmatrix} V_1 \\ n_{11} & n_{01} & n_{10} & n_{00} \end{pmatrix} (S_D P_1 \lambda)^{n_{11}} (S_D (1 - P_1) \lambda)^{n_{01}}$$

$$\times (S_D P_1 (1 - \lambda) + (1 - S_D) \phi)^{n_{10}}$$

$$\times (S_D (1 - P_1) (1 - \lambda) + (1 - S_D) (1 - \phi))^{n_{00}}$$

$$\times \begin{pmatrix} D_1 \\ d_1 & d_0 \end{pmatrix} \phi^{d_1} (1 - \phi)^{d_0}.$$

The reduced model has four parameters and four minimum sufficient statistics.

References

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