

## 3.A3.C Worksheet C – Nested Amplification of Nucleic Acid by PCR for the Confirmation of *Renibacterium salmoninarum*

Case Number \_\_\_\_\_

Date \_\_\_\_\_

PCR Reagent	Lot#	Final Concentration	Stock Concentration**	Volume per Reaction (µL) (to total 50 µL)	Volume for ____ samples
d-H <sub>2</sub> O*		Add to total 49 µL		39.1	
10XBuffer		1X	10X	5.0	
MgCL <sub>2</sub>		1.5 mM	50 mM	1.5	
dNTP's		0.4 mM	10 mM	1	
(+)Primer		20 pMole	20 pMole/µL	1	
(-)Primer		20 pMole	20 pMole/µL	1	
TAQ		2 units/Rx	5U/µL	0.4	
DNA <sup>±</sup> (round 1)		-	-	1 µL	-

\*Add water to Master Mix first, TAQ last. \*\*Change "Stock Concentration" parameters as necessary. Different reagent sources supply varying stock concentrations. †Do not add DNA template until Master Mix reaction tubes have been removed from the reagent mixing (MM) area.

### Primer Sets for *R. salmoninarum* 2<sup>nd</sup> (Nested) Round Amplification

<b>P4 (round 2 forward)</b>	5'-AT TCT TCC ACT TCA ACA GTA CAA GG-3'
<b>M38 (round 2 reverse)</b>	5'-C ATT ATC GTT ACA CCC GAA ACC-3'

Gel Concentration	Weight of agarose (grams)	Volume of Buffer (mL)

### Gel Template (Sample Placement Map)

Ladder Brand / Lot # \_\_\_\_\_

Loading Buffer Brand / Lot # \_\_\_\_\_

Enter sample ID below corresponding well number:

PCR Products (Loaded LEFT to RIGHT)														
<u>1</u>	<u>2</u>	<u>3</u>	<u>4</u>	<u>5</u>	<u>6</u>	<u>7</u>	<u>8</u>	<u>9</u>	<u>10</u>	<u>11</u>	<u>12</u>	<u>13</u>	<u>14</u>	<u>15</u>
<u>16</u>	<u>17</u>	<u>18</u>	<u>19</u>	<u>20</u>	<u>21</u>	<u>22</u>	<u>23</u>	<u>24</u>	<u>25</u>	<u>26</u>	<u>27</u>	<u>28</u>	<u>29</u>	<u>30</u>