## \*Composition Certificate\*

We hereby declare that the composition of the in-vitro diagnostic medical device < Please add product name (without listing codes/catalogue numbers unless needed to identify the product) as it appears in the Declaration of Conformity / CE / Free Sale Certificate / CFG / Canadian Medical Device Active License> is listed in the below table:

< Please fill in the below table according to chosen Raw Material type;</p>
- When "Component" is chosen, please complete the info "Raw Material Component Name" and "Raw Material" and add "N/A" in the rest of columns or delete them as applicable
- When "Active ingredient" or "Inactive ingredient" is chosen as raw material type, please add "N/A" in the column "Raw Material Component Name" or delete it, and complete the info in the rest of the columns >

< If the composition is variable for each variant, please add the composition for each variant as applicable >
< Please clarify any added abbreviations>

Raw Material Type (Please choose either Component, Active ingredient or Inactive ingredient)	Raw Material Component Name (Please add component name of the product where applicable)	Raw Material (Please add raw material name of the listed Component, Active ingredient or Inactive ingredient)	Raw Material Concentration (Please add raw material concentration where applicable)	Raw Material Role (Please add raw material role where applicable)	Raw Material Activity (Please add raw material Activity where applicable)