**Variant of Concern Screening with RT-PACE**

This document contains:

[**Starting method for all sample sizes**](#_For_All_Sample)

[**Protocol for 1536 well plates (robotics)**](#_RT-PACE_with_1536)

[**Protocol for 96 well plates (manual)**](#_RT-PACE_with_Manual)

**Equipment List**

# For All Sample Size Formats

**Starting RNA Materia**

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| Sample quality plays a part in the accuracy of this assay. We have had good results with RNA extracted using the Kingfisher method, but poor results with samples harvested for bacterial studies that were subsequently used for viral RNA. In these instances the results will appear as low and non-specific amplification. Try to limit freeze-thaw prior to genotyping. |

An ideal starting concentration of RNA is 1.5-2.5ng µL-1 or a cT value < 30 in RNase free water.

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| Samples from an RNA extraction will work considerably better than a total nucleic acid extraction. As any RNA measurement will include human as well as viral RNA, this starting amount may vary. If total nucleic acid is used, a considerably higher starting concentration is suggested as the viral RNA will only represent a small amount of the measured concentration. The Nanodrop, which fast and easy, also has a slight over-estimation of RNA concentration.  For clinical samples, an RNA concentration may be difficult to obtain, a cT score from initial testing will suffice. We obtain good results with cT<30 after which the calling becomes more difficult and low or non-specific amplification occurs. |

**Create Primer Set**

Primer sets may be updated as new variants occur. The complete pipeline of PERL scripts along with links to example input data files is available from <https://github.com/pr0kary0te/SARSmarkers>

Primers may be ordered from any oligonucleotide supplier. First ensure that the primer sequences already include the RT-PACE specific tails. Aside from this, the primers require no further modification.

Each SNP marker is probed by a primer triplet which require combining prior to use:

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| --- | --- |
|  | Volume |
| Forward Primer 1 (100µM) | 12µL |
| Forward Primer 2 (100µM) | 12µL |
| Reverse Primer (100µM) | 30µL |

Table 1: Marker master mix. This combination is required for each primer, so a panel of 10 targets will require 30 primers to be ordered and combined into 10 marker master mix as described here.

# RT-PACE with Manual 96-well Capability

**Create the RNA Sample Plate**

Distribute samples into a 96 well plate. 1-2 negative controls (water) and 1-2 positive controls are recommended alongside the testing panel.

For manual testing, a recommended layout is to repeat the same sample across the row of the plate. In this example A1-A12 is the same multiple aliquots of sample 1, B1-B12 is the same multiple aliquots of sample 2, C1-C12 is the same multiple aliquots of sample 1 and so forth. A row of negative control (water) is recommended.

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Figure xx: RNA distribution into sample plate. Each sample replicated 8 times. 5 µL RNA in water in each well.

This layout allows 8 samples to be screened with 12 markers, for different sample/marker combinations different layouts or multiple plates can be used but please adapt the volumes of master-mix solutions accordingly.

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| To prevent contamination during plate filling with PACE mix, the RNA may be dried down within the plate. Spin for 10 seconds then put the plate in an oven/incubator at 48°C for 40 minutes. Cool the plate to room temperature before applying the RT-PACE ready mix.  This step is not essential for the 96-well protocol and may be omitted if required. |

**Apply RT-PACE Mix to the RNA Sample Plate**

Create a master mix of RT-PACE mix and RT-enzyme sufficient for 12 markers. Make immediately prior to use and store on ice. Protect from light.

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|  | Volume |
| RT-PACE mix | 870 µL |
| RT-Enzyme | 8.7 µL |

Table 2: RT-PACE master mix for 12 markers. Scale up or down as required.

Now combine the RT-PACE master mix with other reagents.

Diagram

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|  | Volume | Perform this 12 times for each marker option. |
| RT-PACE master mix from Table 2 | 66 µL |
| RNase-free Water\*\* | - |
| Marker master mix from Table 1 | 5.9 µL |

Table 3: RT-PACE Ready mix. Mix carefully and use immediately. Note that the marker master mix components may separate in storage so ensure it is mixed well before adding to Ready Mix.

\*\* If the RNA was dried in the plate before use, change this to 64 µL and dispense 10 µL per well rather than 5 µL.

Apply 5 µL of the ready mix solution for each marker to the 96 well plate with aliquots of the same marker in columns. In this example A1-H1 is multiple aliquots of marker 1, A2-H2 is multiple aliquots of marker 2 and so forth.

Background pattern

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Figure xx: Marker ready mix distribution into sample plate. Each marker replicated 12 times. 5 µL ready mix added to each well.

Seal carefully using an optically clear seal, laser seal if possible and centrifuge for 10 seconds before PCR. If you are preparing multiple plates before running, store in the fridge protected from light for up to 30 minutes.

**PCR Procedure**

Transfer to thermal cycler, preheated where possible. If plates are not able to be laser sealed, consider hotlid options to reduce volume loss by evaporation.

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| --- | --- | --- | --- |
| Temperature | Time | Cycling |  |
| 50°C | 10 minutes |  |  |
| 94°C | 15 minutes |  |
| 94°C | 20 seconds | 10 cycles | (Dropping 0.8°C per cycle) |
| 65-57°C | 60 seconds |
| 94°C | 20 seconds | 35-40 cycles |  |
| 67°C | 60 seconds |  |

Table 4: PCR parameters. This procedure was optimised using a water bath thermal cycler to reduce ramping time.

After cycling, remove and allow to cool to room temperature (this should only take a few seconds). If laser sealed, spin the plate upside-down for 10 seconds to capture condensation (Only perform this step with laser sealed plates!). With the plate rightside-up, centrifuge for 10 seconds. If condensation persists, allow another 30 seconds of cooling and re-centrifuge.

**Plate Scanning**

Use a scanner fitted with Fl 485/520, Fl 520/560 and Fl 570/610 optic modules. The method of scanning will vary greatly with manufacturer. The aim is to obtain readings for FAM, HEX (same parameters as VIC) and ROX dyes with ROX used for normalisation.

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| While an end-point reaction, it is possible to scan the plate, then return to the thermal cycler for additional cycles before re-scanning. Be careful to save both files not just overwrite. |

**Data Analysis**

Where qPCR is used for scanning, software already exists to create cluster plots and this may be the best choice for lab workflow.

Alternatively use Kraken (may already be owned by DHSC) or KlusterCaller. A free version of Klustercaller exists alongside instruction manual here: <https://www.biosearchtech.com/support/tools/genotyping-software/klustercaller>

Graphical user interface, application

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Graphical user interface

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Using the Klustercaller software, import all the scanned plates, these will automatically appear in the project and will be listed by plate and separated by marker. For each marker, check the cluster plot as an indicator or genotype. In each case the FAM signal will represent one allele and the HEX signal will represent the other allele.

The auto-calling function is very weak, so circle and assign samples to clusters if they cluster well but are not coloured by the software.

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| Be aware that Kraken will adjust the grid axis depending on the samples, so if you only have one allele present, it will auto-scale to make the sample appear heterozygous. Compare axis across the project before submitting for export to ensure that this has not occurred |

Export as “Submitted results” will generate an Excel spreadsheet with each sample compared to each marker.

