

▲ 860µl of prepared sample + 40µl of GPL-2  
▼ 100µl extracted and purified DNA elution  
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▼ 100µl extracted and purified DNA elution

● Lysis Buffer (900µl)  
Columns 1 & 7

● Functionalized Beads (200µl)  
Columns 2 & 8

● Wash #1 Buffer (600µl)  
Columns 3 & 9

● Wash #2 Buffer (600µl)  
Columns 4 & 10

● Elution Buffer (100µl)  
Columns 6 & 12

**QPMKit miQron protocol parameters**

| Step Name     | Column | Volume (µl) | Time (sec) | Mixing Speed (1-10) | Dry Time (sec) | Magnet Capture Time (sec) |
|---------------|--------|-------------|------------|---------------------|----------------|---------------------------|
| Bead Transfer | 2 & 8  | 200         | 30         | 5                   | 0              | 60                        |
| Lysis         | 1 & 7  | 900         | 120        | 5                   | 0              | 60                        |
| Wash #1       | 3 & 9  | 600         | 60         | 5                   | 0              | 60                        |
| Wash #2       | 4 & 10 | 600         | 60         | 5                   | 60             | 60                        |
| Elution       | 6 & 12 | 100         | 300        | 5                   | 0              | 60                        |
| Discard Comb  | 3 & 9  | 600         | 0          | 10                  | 0              | 0                         |

**1** Add 800µl of GPL-1 Buffer to the GPL-2 Buffer tube.  
Ensure the GPL-2 Buffer is mixed well by vortexing.  
! The unused mixture should be kept at 4°C for storage.



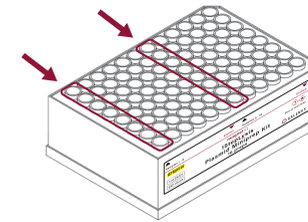
**2** Pellet 1-10ml of bacterial cell sample. Resuspend bacterial pellet in 860µl of Lysis Buffer, then add 40µl of GPL-2 Buffer.  
! For 10ml bacterial culture, add 80µl of GPL-2 Buffer for better lysis.



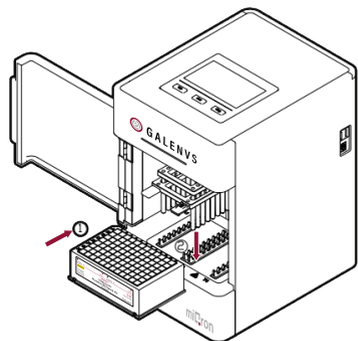
**3** Mix well by pipetting up-down until solution is homogenous.  
Incubate for 5 min at room temperature.



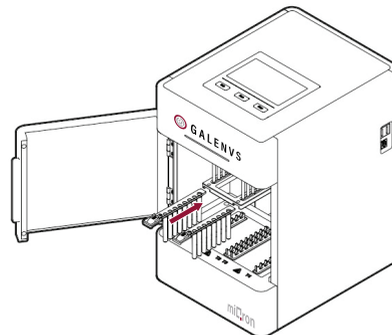
**4** Add the Lysis Buffer (columns 1 and 7).



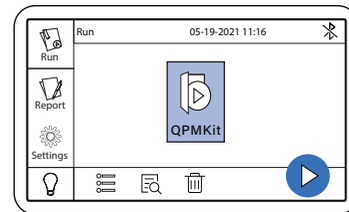
**5** Place plate into the miQron, taking care that the label is facing outward.



**6** Insert two combs.

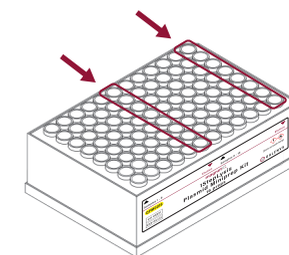


**7** Select the QPMKit protocol and press play.



When program is complete, remove plate from miQron and discard combs.

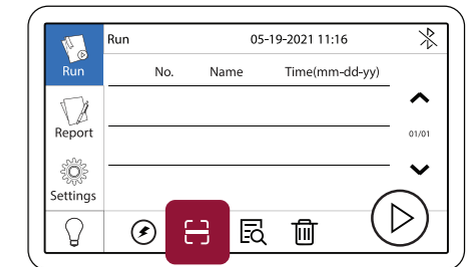
Columns 6 and 12 contain the purified DNA elution.



**miQron**

**Protocol Update**

To import the updated protocol into the miQron, press the **Scan Protocol** icon from the **Run** menu (protocol list view window). Use the scanner on the QR code below.



QPMKIT PROTOCOL V1.0