# **Streptococcus pyogenes Detection with real time PCR reagents**

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S. pyogenes FAM-BHQ1	PP2800	0.055ml	2				
AttoMaster 2X Mix for qPCR	AM10	1.25 ml	2				
S. pyogenes Plasmid 200 pg/ml	PLAS2800	0.25ml	2				
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•	S. pyogenes FAM-BHQ1 PP2800 0.055ml AttoMaster 2X Mix for qPCR AM10 1.25 ml S. pyogenes Plasmid 200 pg/ml PLAS2800 0.25ml						
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### Overview:

Ref. 2

Streptococcus pyogenes, Group A strep, causes the common tonsillitis, pharyngitis and impetigo. Group A strep is also the cause of serious, life-threatening, cellulitis and systemic disease. Immune processes are responsible for post-streptococcal rheumatic fever and glomerulonephritis.

Primers and probe are available for amplification and detection of Group A strep. The Group A strep primers are specific for a 98 bp

portion of the SpecB gene. The Group A strep detection probe is a molecular beacon (1)

### **Products**

Products Catalog No. Quantity

S. pyogenes FAM-BHQ1 PP2800 0.055ml

Store at -20C. 25- 20µl reactions

Typical use: Add 2X master mix and DNA, then thermal cycle.

Attostar reagent contains primers and probe.

Detection at 510nm.

AttoMaster 2X Mix for qPCR AM10 1.25 ml

Store at -20°C. 125-20 ul reactions

Contains Taq polymerase (requires heat activation), dNTPs (0.4 mM) with optimal dUTP to dTTP ratio, heat labile UDG, Mg(6 mM), and buffer.

Typical use: Add Attostar Primer-probe, DNA, and then thermal cycle.

S. pyogenes Plasmid 200 pg/ml PLAS2800 0.25ml

Store at –20C.

Typical use: make serial 10 fold dilutions in TE for standard curve, diluting 5 ul into 45 ul TE buffer.

## **Detection of S.pyogenes DNA:**

# Thermal cycle conditions for PCR reactions on RotorGene\*

\*Similar cycle conditions and reaction volumes may be used on many other thermal cyclers.

```
25°C 10 min (UDG treatment time)
95°C 120 sec (activation for AttoMaster polymerase)
40 cycles
95°C 15 seconds
60°C 30 seconds RotorGene Channel Setup FAM/Sybr, Cy5; Gain 7
72°C 30 seconds
```

FAM/Sybr has a source of 470nm and Detector 510nm (LightCycler use F1) Cy5 has a source of 625nm and Detector 660hp nm Quasar 670 has the same fluorescent absorption and emission as Cy5.

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# Thermal cycle conditions for PCR reactions on LightCycler

```
25°C 10 min (UDG treatment time)
95°C 120 sec (activation for AttoMaster polymerase)
40 cycles
95°C 15 seconds
60°C 30 seconds acquire fluorescent signal on F1 gain =1
72°C 30 seconds
40°C 30 seconds cool
```

# Use of the Streptococcus pyogenes plasmid:

Dilute the plasmid in TE to prepare a standard curve. Common dilutions would be 10-fold from 200 to 0.002pg/ml. The 0.02~pg/ml plasmid dilution contains 12 copies of plasmid in 2  $\mu l$ .

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# Attostar Primers/probe PP2800 RotorGene-PCR 20µl reactions

	RotorGene Reagents needed for 20 ul PCR final reaction tube volumes										
Reaction tube number	1	2	3	4	5	6	7	8	9	10	
Attostar Primer-Probe											
FAM labeled (10X)	2	4	6	8	10	12	14	16	18	20	μl
Master mix (2X)	10	20	30	40	50	60	70	80	90	100	μl
Dispense 12 ul / reaction tube											
			Add	8 ul DNA /	reaction tub	е					1

# Attostar Primers/probe PP2800 LightCycler-PCR 20 $\mu$ l reactions

	LightCycler Reagents needed for 20 ul PCR final reaction tube volumes											
Reaction tube number	1	2	3	4	5	6	7	8	9	10	1	
Attostar Primer-Probe											П	
(10X)	2	4	6	8	10	12	14	16	18	20	μ	
Master mix (2X)	10	20	30	40	50	60	70	80	90	100	þ	
BSA 1 mg/ml	1	2	3	4	5	6	7	8	9	10	μ	
Dispense 13 ul / reaction tube										Ī		
			Add	7 ul DNA /	reaction tub	е						

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# Extraction / amplification control with T4 bacteriophage

Adding T4 bacteriophage (BAC130) to the sample provides DNA for extraction, amplification, and reaction condition PCR controls.

When added to a sample, T4 adds a known amount of DNA. The T4 DNA can then be extracted, amplified, and detected as a control. T4 controls for the efficiency of DNA extraction, the presence of PCR amplification inhibitors, intact amplification reagents (DNA polymerase, buffer, dNTPs), and instrument function (thermal cycling and fluorescent detection system).

The T4 DNA may be detected in a separate PCR reaction (using FAM labeled T4 probe in PP100). Or the T4 DNA and test organism DNA may be detected using a multiplex reaction (using Quasar 670 labeled T4 probe in PP160 and FAM labeled test organism probe).

# Brief procedure for use of T4 as extraction and amplification control:

O Add 5μl T4 bacteriophage to the sample. Proceed with DNA extraction. Dilutions of the bacteriophage may be made to give a final PCR Ct value that is about 35. At this dilution, the phage is more sensitive, i.e. more likely, to detect a poor extraction or the presence of PCR inhibitors in the reaction.

Please refer to the BAC130, PP100, and PP160 product literature.

# T4 multiplex reaction (PP160)

# RotorGene Multiplex PCR 20µl reactions

	Multiplex Reagents needed for 20 ul PCR final reaction tube volumes										
Reaction tube number	1	2	3	4	5	6	7	8	9	10	)
Attostar Primer-Probe PP160	2	4	6	8	10	12	14	16	18	20	μl
Primer-Probe mix for TEST organism	2	4	6	8	10	12	14	16	18	20	μl
2 x master mix	10	20	30	40	50	60	70	80	90	100	μl
Dispense 14 ul / reaction tube											
			Add	d 6 ul DNA	reaction tu	be					

Reaction tube number	11	12	13	14	15	16	17	18	19	20	)
Attostar Primer-Probe PP160	22	24	26	28	30	32	34	36	38	40	μl
Primer-Probe mix for TEST organism	22	24	26	28	30	32	34	36	38	40	μl
2 x master mix	110	120	130	140	150	160	170	180	190	200	μl
Dispense 14 ul / reaction tube											
	Add 6 ul DNA / reaction tube										

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#### Reference:

1) <a href="http://www.molecular-beacons.org/Introduction.html">http://www.molecular-beacons.org/Introduction.html</a>

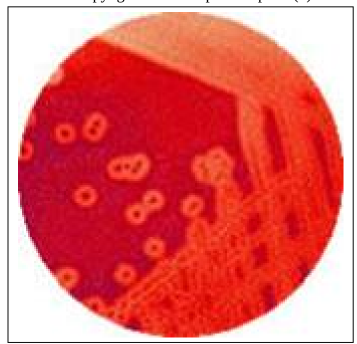
2) Image:S. pyogenes beta hemolytic colonies: <a href="http://www.jnu.ac.in/Faculty/ajohri/">http://www.jnu.ac.in/Faculty/ajohri/</a>

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# **Enlarged image:**

Beta hemolytic colonies of S. pyogenes on sheep blood plate (2).



10-18-07