

Date 2/14/2013

Objective PCR amplification of triplet repeat region of FMR1 with a modified program (as below):

- Without hot start
- Three step amplification process
- Denat temp set to 92degC/ 30secs
- Annealing temp set to 57.8deg/ 1:30mins
- Extension temp. grad (55-75deg); time set to 2:30mins

Description *Template used:* 100ng genomic DNA NA20230, (53/54 rpt sample from Coriell)
Primers: FT for and FT rev
Polymerase: Native Taq Polymerase (Life Technologies)
Solvents: No Solvt rxns
 Rxns in 0.5M NMP + 2.2M Betaine

Reaction Composition:

Component	0.5M NMP + 2.2M Betaine rxns Final Conc/ amt
Taq Buffer	1 X
MgCl ₂	1.5 mM
dNTPs	0.4 mM
FT-For/ SFS-for	0.4 uM
FT-Rev/ SFS-rev	0.4 uM
gDNA NA203230	100 ng
Taq Polymerase	2.5 U
Water	
NMP	0.5 M
Betaine	2.2 M
total	25ul

Cycling Conditions:

- 1 92degC/ 30secs
- 2 57.8degC/ 1:30mins
- 3 55-75degC/ 2:30mins
- 4 GOTO 2 40 times

Gradient steps:

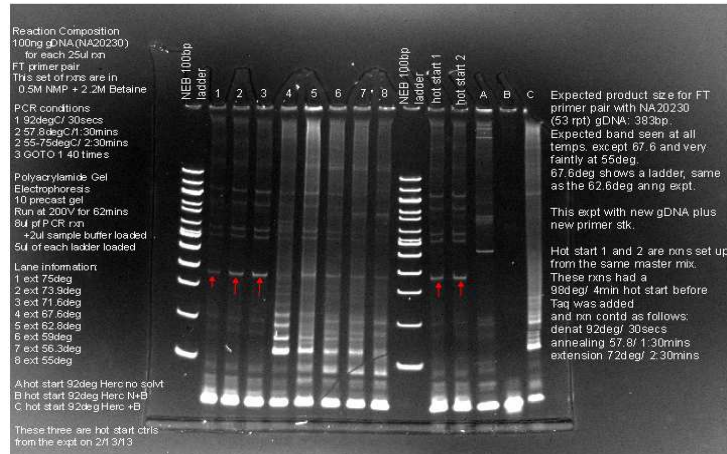
75, 73.9, 71.6, 67.6, 62.8, 59, 56.3, 55degC

Gel Electrophoresis:

10% precast polyacrylamide gels (Life Technologies)
 8ul of PCR rxn + 2ul of (5X) sample buffer loaded
 Molecular Ladders: 5ul loaded
 100bp ladder (NEB)
 50bp ladder (NEB)

Gel Pictures

Lanes are marked according to the ext temp of the particular rxn
Please ignore Lanes marked A, B, C (not relevant to this expt)
FT primers - NMP + Betaine (20sec exposure)



Comments Reactions have worked at all temps 75-55 but look best at 72.
Extension 67.6 has a ladder (also seen with anng 62.6 ext 67.6,
see report ext 55-75 anng 62.6)