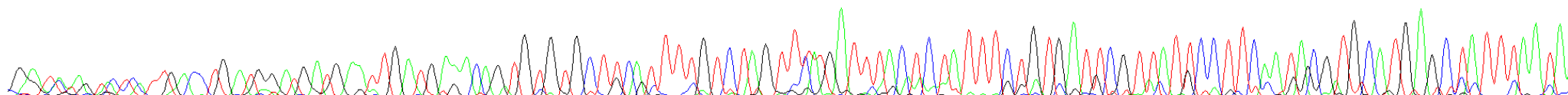


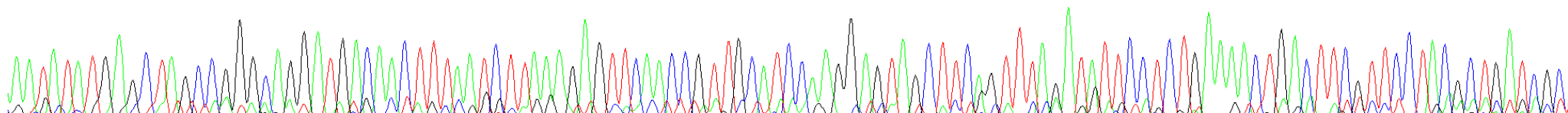
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Sample: 3g54470F68_3g54470F Lane: 19 Base spacing: 14.03 909 bases in 10929 scans Page 1 of 2

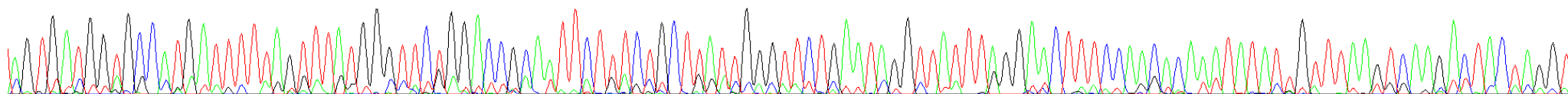
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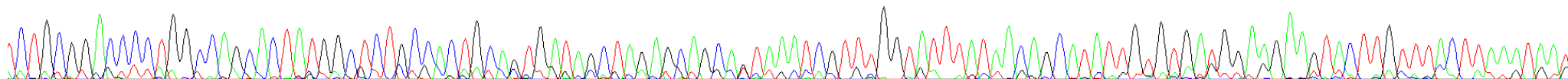
130 140 150 160 170 180 190 200 210 220 230 240 250
A A T A T A T G A G C T A G C C G G G C A G G A T G A C A A C T T T A A T C T T A A A G A G T T C A A C C G T T G C A T C C A A G G A G T A G C T T C A G T T T A G A T A T T C C T C T G A A A A C T G A C T T C G T T C C T A C G C T G A T C G C T



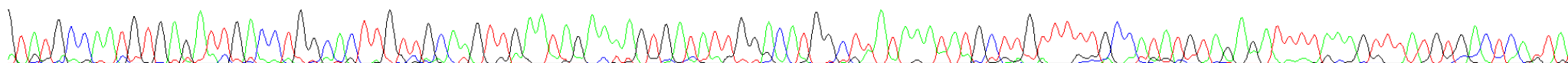
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A G T G A T G G T G C C A T G A T T T T T A G T T T A T G G G T T C T G G A C C G C A A T T C T T T C T G C T T A T T G G G T T C T G A A T A G G T T A T T T A G G A A C T T T C C A A C A C A A A T A T A T T G T T T A A G T C A A G A C T A C T A G G T



380 390 400 410 420 430 440 450 460 470 480 490 500
T C T G C G G A C C C C T G G C C A G C A C T A T G G C T C T G C C A C T G C A G T G A T A G C T A G A G C A G C A G T A A A T C G T T T G G T A T T T T A A C A G C A T A T T G T G T G A T G G A A G A A G T A C T T G T T T A C T T A A A T A A T



510 520 530 540 550 560 570 580 590 600 610 620
G T A T G C C A A T G T G A G A T T G A C C T G G C A C T T G T T G C A A G T T G A A T A G A A A A G T G A T A T T G G A C A T G G C T T A T A A A T A T G C T T G T T T T T G C C A T A T G T A G A G A T T T T A A A G T T T A T G T G A C T C A T T A T



File: 050516-09_N06_3g54470F68-3g54470F.ab1 Run Ended: 2005/5/18 10:9:39 Signal G:204 A:209 C:168 T:293
Sample: 3g54470F68_3g54470F Lane: 19 Base spacing: 14.03 909 bases in 10929 scans Page 2 of 2

630 640 650 660 670 680 690 700 710 720 730 740 750
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760 770 780 790 800 810 820 830 840 850 860 870
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880 890 900
TG GTGT G TG G TTTG TG G TTT TGGTGTATTTT G

