Genomic DNA extraction from single fly

1- Set hybridization oven at 55°C
2- Make sure HB buffer is not precipitated. If it is - warm it up until precipitate dissolves and mix before use
3- Homogenize single fly with 250 ul of HB buffer (see recipe at the end of this protocol)
4- Add 5 ul of 10 mg/ml of Proteinase K to each tube, mix
5- Incubate at 55°C 2-3 hr
6- Add 85 ul of 5 M NaCl and vortex
7- Spin down pellet for 20 min at full speed
8- Transfer supernatant into new tubes
9- Add 340 ul of 100% cold EtOH, mix, put on ice for 10 min, spin for 20 min at full speed in the cold room to pellet DNA
10- Remove EtOH with blue tips without disturbing pellets
11- Add 1 ml of 70% EtOH, invert few times and spin for 15 min at full speed
12- Remove EtOH using blue tips
13- Quick spin tubes
14- Remove all leftovers of EtOH with yellow tips without disturbing pellets
15- Dry DNA until you don’t see any liquid in the tube
16- Add 100 ul of water
17- Leave o/n in refrigerator
18- Next day: flick and spin. Freeze samples on –20 C until needed

HB buffer (olga’s bench): 50 mM Tris pH 7.5
400 mM NaCl
20 mM EDTA
0.5% SDS
to make HB combine:
39 ml of water
2.5 ml of 1M Tris pH7.5
4 ml of 5M NaCl
2 ml of 0.5 M EDTA pH 8.0
2.5 ml of 10% SDS