Notes on the fly project

J. Ignacio Lucas Lledó
October 24, 2016

1 Delivery of oligonucleotides

The oligonucleotides designed on 2016-06-09 were ordered to biomers.net, and delivered. They are dry. Table 1 specifies the yield and the corresponding amount of buffer required to dilute them to 100 pmol/ μ l.

2 DNA extraction test

On September 13th 2009, Dr. Carazo, Mrs. Sultanova, Mr. Andiç and I started the DNA extraction from 8 flies, following the GBS protocol. Table 2 summarizes the procedure. Not that we needed 4 people to run this, but they all were interested in going through the protocol, which is great. The protocol is applied to each fly individually, with the aim of measuring how much DNA we can get from one fly. Four of the flies had their oviduct previously removed by Mrs. Sultanova. We want to make sure that the ovariectomy does not significantly reduce DNA yield.

We ended up measuring DNA concentration 3 times, due to problems with the Qubit standards. The error 'Incorrect standards' was apparently caused by the fact that the reagent had been kept in the fridge. I thought that it would be enough to wait for it to thaw, to bring it to room temperature. Plus, it is a very small amount of it that gets dissolved in the buffer, at room temperature. For whatever reason, the kit worked once we kept the reagent at room temperature. I thank Rebeca Dominguez, from the Evolutionary Genetics lab, her help to solve the problem.

The slight difference in average DNA concentration between control and ovariectomy samples is not significant, which is good news. I expect three samples, namely 1, 3, and 6, to have enough DNA left to run the rest of the protocol with them. I could add sample 5 to see what we get from it.

Table 1: Name and available amount (yield) of oligonucleotides received from biomers.net. 'Volume' makes reference to the volume required to dissolve the corresponding oligonucleotides at 100 pmol/ μ l (μ M).

Name	Yield	Volume	Name	Yield	Volume
	(nmol)	$(\mu \mathrm{l})$		(nmol)	$(\mu \mathrm{l})$
Top1.1	82.6	826	Bot1.1	64.8	648
Top1.2	106.8	1068	Bot1.2	110.4	1104
Top1.3	94.9	949	Bot1.3	92.6	926
Top1.4	85.6	856	Bot1.4	75.8	758
Top 2.1	78.9	789	Bot2.1	90.7	907
Top 2.2	81.9	819	Bot2.2	97.4	974
Top 2.3	100.5	1005	Bot2.3	84.2	842
Top2.4	88.4	884	Bot2.4	82.3	823
Top3.1	98.6	986	Bot3.1	87.1	871
Top3.2	95.2	952	Bot3.2	96.6	966
Top3.3	100.2	1002	Bot3.3	87.9	879
Top3.4	95.6	956	Bot3.4	78.7	787
A1.1	60.2	602	A2	65.9	659
A1.2	52.6	526	R1	78.7	787
A1.3	56.7	566	R2	60.0	600
A1.4	51.0	510			

Table 2: Summary of the DNA extraction test.

Sample	Treatment	Elution (μl)	$\operatorname{Conc.}(\operatorname{ng}/\mu\operatorname{l})$	Yield (ng)
1	_	100	4.240	424
2	_	100	0.978	98
3	_	100	3.600	360
4	_	100	1.790	179
5	ovariectomy	100	1.990	199
6	ovariectomy	100	3.980	398
7	ovariectomy	100	1.140	114
8	ovariectomy	100	1.530	153

3 Literature review

Original motivation of classic life span QTL mapping studies was the elucidation of metabolic pathways involved in aging [12]. I suspect that the biomedical interest in the mechanisms of aging preceded the interest in naturally occurring variation, as well as the real effect of the QTL identified in natural conditions. Several sex-specific life span QTL exist in autosomes of D. melanogaster [13, 15]. These QTL were originally identified using recombinant inbred lines (RIL) descended from two strains [13]. In this design, the dominance of the QTL is not known, since only homozygous genotypes are produced [11, page 432]. However, the quantitative complementation test with deficencies used to refine the QTL [15] seems to assume that one of the alleles is at least partially recessive.

Crow [5] mentions the genetic basis of heterosis as one of the main midcentury controversies in population genetics. He is very clear that the debate between dominance and overdominance was resolved in favor of the dominance hypothesis. That is, both the inbreeding depression and the hybrid vigor (heterosis) are mostly due to recessive deleterious alleles that become either expressed upon inbreeding or concealed upon hybridization. A review [3] cites another work by James F. Crow that I cannot access (volume 9 of the Oxford Series in Evolutionary Biology), in which the author shows evidence of 30% of autosomes isolated from natural populations of *D. melanogaster* being lethal in homozygosis. The same cannot be true for sex chromosomes, of course. Among the 70% of natural chromosomes that are not recessive lethals, homozygosity reduces survival to adulthood in 84%.

Full-sib mating during 10 generations is expected to produce an inbreeding coefficient of 0.886. This is the probability at any locus that the two copies inherited from the parents are identical by descent; that is, they are exact copies of the same chromosome in one of the recent ancestors. But how is this identity-by-descent distributed along a chromosome? It is not random, but clustered, because of linkage. We expect an alternative sequence of fragments that are either identical by descent or not. That is, the inbreeding itself already produces a mosaic of heterozygosity and homozygosity. The question is how long the pieces are. Too short IBD tracts would require very dense markers. According to I. R. Franklin [6], previous authors (Bennet and Fisher) had supposed that 'after a long number of generations of inbreeding the number of heterogenic segments will be approximately distributed as a Poisson variable, and that the distribution of the length of each segment (x)will tend to a negative exponential $(1/a)e^{-x/a}$. However, this assumptions are not satisfactory, especially for the first few generations of inbreeding. Taking into account that there is no crossing-over in *Drosophila males*, the

Table 3: Digestion reactions.

Sample	$\begin{array}{c} \text{DNA} \\ (\mu \text{l}) \end{array}$	H_2O (μl)	$10 \times \text{ buffer} $ (μl)	$\begin{array}{c} \text{NspI} \\ (\mu l) \end{array}$	Total (μl)
1	80	207	32	1	320
3	80	207	32	1	320
5	80	207	32	1	320
6	80	207	32	1	320

probability that two loci in the same chromosome are homozygous by descent after one generation of sib mating is $(1/32)(3+3\lambda_f+\lambda_f^2+\lambda_f^3)$ [6], where λ_f is the linkage parameter $1-2y_f$, representing the probability of recombination in the female.

The suggestion given by Dr. Carazo of using simulations is actually good. I started learning to use simuPOP.

4 Enzymatic digestion of DNA samples 1, 3, 5, and 6

I run the digestions as described on table 4, for above 4 hours, at 37°C. Together with Muhammed Andiç, we also cleaned up the reaction with magnetic beads. We used a 1:1 ratio of beads to DNA, and eluted with water (40 μ l).

5 Literature review

From page 489 of [16]: "The adaptive significance of achiasmatic meiosis is not entirely clear. Obviously it has been evolved repeatedly in at least ten major groups of animals. In those cases where achiasmatic meiosis is confined to one sex, we have far too little information as to the chiasma frequency in the other sex. If it were generally true that in the 'chiasmatic sex' there is only one chiasma per bivalent, we might assume that selection for a low level of genetic recombination per se had led to total loss of chiasmata in one sex. If, on the other hand, absence of chiasmata in one sex is generally accompanied by a fairly high chiasma frequency in the other, this explanation obviously would not hold. Only evidence from species that had acquired achiasmatic mechanisms fairly recently in their phylogeny would be significant from this

point of view (*Tipula caesia* would be relevant – *Drosophila* spp. would not). The only evidence on this point seems to be the grasshopper *Thericles whitei*, with no chiasmata in the male and a very low chiasma frequency in the female; it obviously supports the hypothesis of selection for low levels of recombination."

However, selection for low levels of recombination would not explain why the achiasmatic sex is nearly always the heterogametic sex [2]. White also mentions the hypothesis of achiamatic spermatogenesis being an adaptation to tolerate inversion-heterozygosity. However, no evidence supports this hypothesis.

More recently, [10] mentions two more hypotheses: first, that achiasmy may have evolved as a side effect of selection to suppress recombination between the sex chromosomes [7, 8]. And second, that it evolved as a way to promote tight linkage without suppressing recombination on the X or Z chromosome [9].

Well, I cannot review now all the hypotheses now. If anything, I should relate the results of the simulations to the theoretical expectations of the variance in inbreeding coefficient [4, 6]. It seems that multiple linked loci in sex chromosomes have received relatively little attention in the literature [1, 14].

6 Annealing of adapters

On September 28th, Muhammed and I prepared 100 ml of 1× TE buffer, following a recipe online: 1.0 ml 1M Tris pH 8.0, 0.2 ml EDTA 0.5M, and 98.8 ml $\rm H_2O$. We also prepared annealing buffer, according to the protocol, and suspended all the oligonucleotides in TE buffer to 100 μ M, according to the volume specified on table 1. Finally, we prepared 10 μ l of annealed adapters of each type.

7 Quantification with Agilent Bioanalyser

Amparo Martínez, from the Sequencing Service, sent the results from the Bioanalyser run. The four samples have a very similar profile (Figure 1), which looks like the result of a successful digestion followed by a more aggressive selection of long fragments that I expected from a 1:1 ratio of magnetic beads to DNA. Next time, I should use a more generous proportion of beads. Table 4 shows some additional details. I note that the amount of DNA available is now an order of magnitude lower than what I had for the *Coregonus* samples.

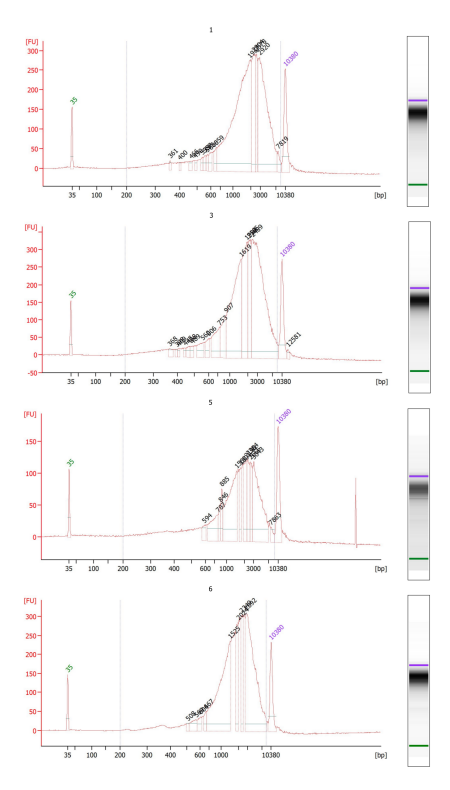


Figure 1: Electropherograms from the high-sensitivity DNA assay of the four samples.

Table 4: Distribution of DNA fragments between 40 and 9000 bp. 'Size' refers to the average fragment size. For total mass of DNA left over, a volume of 35 μ l is assumed to be left. The average fragment mass (Frag.Mass) is calculated in two ways. First, by dividing the overall concentration by the overall molarity (Frag.Mass.1); and then, by applying the formula for the molecular mass of double stranded DNA to the average length of the fragments (i.e., 607.4x + 157.9 g/mol; Frag.Mass.2).

Sample	Size (bp)	Conc. $(pg/\mu l)$	Molarity (pmol/l)	Left over (ng)	$\begin{array}{c} {\rm Frag.Mass.1} \\ {\rm (ng/fmol)} \end{array}$	$\frac{\rm Frag. Mass. 2}{\rm (ng/fmol)}$
1	2206	1425.92	3042.0	49.91	0.468744	1.34008
3	2239	1439.86	2484.9	50.40	0.579444	1.36013
5	2022	1113.97	3509.8	38.99	0.317388	1.22832
6	2376	1523.30	2066.7	53.32	0.737069	1.44334

8 Ligation of adapters

In folder 2016-10-04, I calculate the composition of the ligation reactions to have a 10-fold excess of adapters to DNA fragment ends. On table 5 I show the composition and final concentrations of the working solutions of the four adapters, required to prepare the ligation reactions specified on table 6. Importantly, I decide to assign adapters 1.1, 1.2, 1.3, and 1.4 to samples 1, 3, 5, and 6, respectively. See table 7.

Just to have a graphic representation of the adapters and the expected

Table 5: Composition and final concentration of the working solutions of adapters for the ligation reactions on table 6. The buffer must be $1\times$ annealing buffer. Amounts are μ l, except the final molarity, in the last column, in pmol/ μ l.

Sample	Adapter	$1 \times $ buffer	15 $\mu\mathrm{M}$ Stock	Total	$\mathrm{pmol}/\mu\mathrm{l}$
1	1.1	8.580	1.420	10.00	2.13
3	1.2	8.840	1.160	10.00	1.74
5	1.3	8.362	1.638	10.00	2.46
6	1.4	9.036	0.964	10.00	1.45

Table 6: Volumes of reactants in the ligation reactions. All in μ l. T4 ligase is at 2000000 U/ml.

Sample	DNA	$10 \times$ Buffer	Adapter	1.5 M NaCl	T4	${\rm H_2O}$	Total
1	35	4.40	1.00	1.10	2.00	0.50	44.00
3	35	4.40	1.00	1.10	2.00	0.50	44.00
5	35	4.40	1.00	1.10	2.00	0.50	44.00
6	35	4.40	1.00	1.10	2.00	0.50	44.00

Table 7: Adapters and codewords assigned to the DNA samples.

Sample	Adapter	Codeword
1	1.1	TTGATCCAGT
3	1.2	GATCAGGCAGT
5	1.3	CCAGCTTGT
6	1.4	AGCTGAAT

DNA fragments, this is how they should look like after ligation (adapter 1.1 represented):

The codeword is in red, and the genomic fragment in blue. Note that the restriction site, RCATGY, is not present in the ligation product (R is G or A, and Y is C or T). This allows me to digest a second time, and get rid of chimeric fragments.

On October 5th, I prepared the working solution of the annealed adapters, and the following day I set up the ligation reaction. I then realized the available T4 DNA ligase was at 400000 U/ml, instead of 2000000 U/ml. I decided to double the amount of T4 ligase, and adjust the amount of buffer. I also noticed that sample 6 did not have more than 30 μ l of DNA. Thus, I changed the composition of the reaction accordingly. However, I note that I used 1 μ l of adapters at a slightly higher concentration than necessary to achive the 10-fold excess, because the working solution of adapters was already prepared, and I didn't want to re-calculate the whole thing. See table 8 for the current composition of the ligation reaction.

Table 8: Current composition of the ligation reaction, after necessary adjustments. All amounts are in μ l. T4 ligase was actually at 400000 U/ml.

Sample	DNA	$10 \times$ Buffer	Adapter	1.5 M NaCl	T4	${\rm H_2O}$	Total
1	35	4.62	1.00	1.10	4.00	0.50	46.22
3	35	4.62	1.00	1.10	4.00	0.50	46.22
5	35	4.62	1.00	1.10	4.00	0.50	46.22
6	30	4.02	1.00	1.10	4.00	0.25	40.37

Table 9: Second digestion reactions.

Sample	$\begin{array}{c} {\rm DNA} \\ (\mu {\rm l}) \end{array}$	H_2O (μl)	$10 \times \text{ buffer}$ (μl)	$\begin{array}{c} \mathrm{NspI} \\ (\mu \mathrm{l}) \end{array}$	Total (μl)
1	46.22	119.17	18.49	1	184.88
3	46.22	119.17	18.49	1	184.88
5	46.22	119.17	18.49	1	184.88
6	40.37	103.96	16.15	1	161.48

9 Second digestion

The second digestion is meant to break chimeric fragments. The adapters will not be cleaved because they were designed not to reproduce the cut site. I do not want to clean up the ligation reaction. In the worse case, I will not get rid of the chimeras. But I don't want to loose DNA. So, I count the total ligation volumes (table 8) as DNA volumes in the second digestion reactions specified on table 9.

10 Size selection

I plotted the expected shape of the electropherogram, using data from the in silico digestion with the pattern RCATGY (2016-04-12). The result, shown in figure 2, is remarkably similar to the observed electropherograms after digestion (figure 1)

In any case, from those figures it is not immediate to know the number of loci in a size range. The use of magnetic beads for size selection is tricky, because very precise volumes need to be measured, some DNA is lost, and the range is wider and less reproducible than if selected with a Blue Pippin.

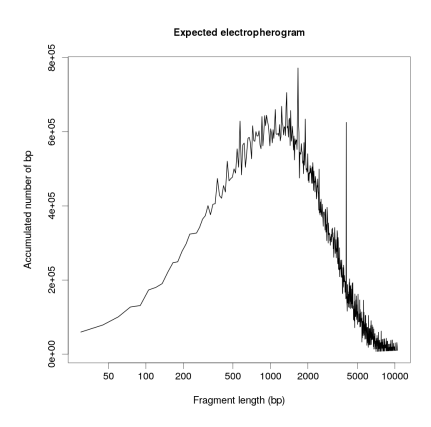


Figure 2: Electropherograms from the high-sensitivity DNA assay of the four samples.

Table 10: Volumes of DNA sample, bead suspension and ethanol to be added during right-side selection (Beads 1, 0.600:1 ratio), left-side selection (Beads 2; 0.645:1 ratio), and washing steps (ethanol). All in μ l.

Samples	DNA	Beads 1	Beads 2	Ethanol
1, 3, 5	184.88	110.93	8.32	2×305.00
6	161.48	96.89	7.27	2×305.00

Table 11: Amplification primers

	Primer 1a	Primer 1b	Primer 1c	Primer 1d	Primer 2
Length (bp)	63	63	63	63	 55
Molecular weight (g/mol)	19511.7	19545.7	19536.7	19536.7	16719.9
Basic T_M First cycle (°C)	65.7	65.7	65.7	65.7	64.4
$52 \text{ mM Na}^+ \text{ T}_M 1^{st} \text{ cycle (°C)}$	75.0	75.0	75.0	75.0	73.3
Basic T_M Later cycles (°C)	75.7	76.4	76.4	76.4	74.3
$52~\mathrm{mM~Na^+~T}_M~\mathrm{Later~(^{\circ}C)}$	87.2	88.0	88.0	88.0	85.7

So, wide size ranges are convenient in this case. Therefore, I should move the range in the direction of lower number of fragments, so that a tighter control in the number of fragments is possible. After looking at the ratios of beads to DNA required for each range, I realize I cannot aim at fragments just between 500 and 800 bp, but rather between 450 and 1000 bp. Because I will use 300 bp, paired end reads, I stick to this range. Finally, I use beads to sample ratios of 0.600 (right-side selection at \sim 900 bp) and 0.645 (left-side selection at \sim 350 bp. I expect less than 33000 loci, according to results from the 2016-04-12/RCATGY folder. Given that the total volumes are either 184.88 or 161.48 μ l, this translates into the volumes specified on table 10. I will attempt elution with 30 μ l or less.

11 Amplification PCR

After size selection I do not expect to have more than 2 ng of DNA per sample. I will not quantify before amplification, in order to save DNA and because I would not trust so small measures.

Table 12: PCR reactions in 50 μ l. 10 μ l of 10 μ M Primer Mix may be prepared with 1 μ l 100 μ M Primer 1 + 1 μ l 100 μ M Primer 2 + 8 μ l nuclease-free water.

Component	Volume (μl)	Final conc. (pmols/l)
$10 \ \mu M$ Primer Mix	1.0	200000.00
Template DNA	15.2	580.24
$2 \times$ Phusion Master Mix	25.0	_
Nuclease free water	8.8	_
Total	50.0	

12 Qubit quantification and pooling

13 Bioanalyser quantification of pooled library

14 Sequencing

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