Report: Hybrid vigor in response to Eimeria in the HMHZ

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To be fixed before all
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To be fixed before all

• Some information regarding latitude and longitude are missing for the following mice:

SK 3174

• We still miss info (HI) on the following mice (ask Jarda):

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AA_0411, AA_0412, AA_0420, AA_0464, SK_2668, SK_2669, SK_2671, SK_2674, SK_2675, SK_2676, SK_2677, SK_2677, SK_2678, SK_2681, SK_2682, SK_2684, SK_2685, SK_2687, SK_2688, SK_2690, SK_2692, SK_2693, SK_2695, SK_2696, SK_2699, SK_2700, SK_2701, SK_2702, SK_2703, SK_2704, SK_2705, SK_2710, SK_2713, SK_2715, SK_2724, SK_2727, SK_2729, SK_2733, SK_2734, SK_2736, SK_2737, SK_2738, SK_2739, SK_2745, SK_2750, SK_2751, SK_2751, SK_2752, SK_2754, SK_2755, SK_2756, SK_2758, SK_2759, SK_2760, SK_2761, SK_2775, SK_2778, SK_2780, SK_2782, SK_2789, SK_2792, SK_2793, SK_2794, SK_2795, SK_2795, SK_2799, SK_2800, SK_2801, SK_2802, SK_2803, SK_2804, SK_2805, SK_2851, SK_2852, SK_2853, SK_2854, SK_2855, SK_2856, SK_2857, SK_2858, SK_2859, SK_2860, SK_2861, SK_2862, SK_2863, SK_2864, SK_2865, SK_2866, SK_2868, SK_2869, SK_2871, SK_2873, SK_2874, SK_2875, SK_2876, SK_2877, SK_2878, SK_2879, SK_2880, SK_2881, SK_2884, SK_2885, SK_2886, SK_2886, SK_2886, SK_2886, SK_2881, SK_2884, SK_2885, SK_2886, SK_2886, SK_2887, SK_2884, SK_2885, SK_2886, SK_2886, SK_2886, SK_2881, SK_2884, SK_2885, SK_2886, SK_2886, SK_2886, SK_2881, SK_2884, SK_2885, SK_2886, SK_2886, SK_2887, SK_2884, SK_2885, SK_2886, SK_2887, SK_2888, SK_2889, SK_2958, SK_2959, SK_2960, SK_2961, SK_3174, SK_3333, SK_3417, SK_3461, SK_3462, SK_3469, SK_3470, SK_3471
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General informations on HMHZ

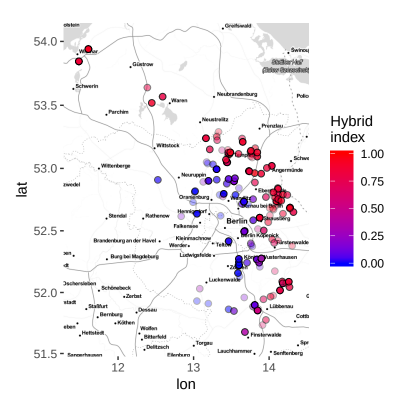


Figure 1: Map of the mice with OPG, PCR or qPCR status, caught in the Brandenburg-MVP transect in 2015, 2016 and 2017. Each point corresponds to one location, a less pronounced transparency indicating more animals sampled at this location. Hybrid index is represented by a gradient from blue (M.m.d) to red (M.m.m)

- 655 mice were captured over three years, from 157 farms
- From these mice:
- 485 mice had Eimeria detected by feces flotation,
- 653 mice had Eimeria detected by colon content PCR (cf paper Victor),
- 160 mice had Eimeria detected by qPCR on intestinal tissues
- On average, 4.04 mice were caught per farm (95% CI 0.34)
- Hybrid indexes were calculated as ratio of M.m.d/M.m.m alleles (between 4 and 14, on average 13 loci)

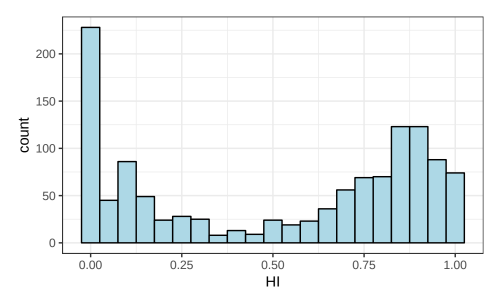


Figure 2: Number of animals caught along the hybrid index

Comparison of prevalences based on detection method

Table 1: Prevalence of Eimeria per year, based on oocyst flotation

	2014	2015	2016	2017
FALSE	0	92.0	126	167.00
TRUE	0	10.0	24	66.00
$\operatorname{prevalence}(\%)$	NaN	9.8	16	28.33

Table 2: Prevalence of Eimeria per year, based on PCR detection. A mouse was considered infected by Eimeria ifone of the 3 markers (COI, 18S or ORF470) gave a sequence

	2014	2015	2016	2017
negative	53.00	110.00	146.00	226.0
positive	23.00	12.00	20.00	63.0
$\operatorname{prevalence}(\%)$	30.26	9.84	12.05	21.8

Table 3: Prevalence of Eimeria per year, based on qPCR

	2014	2015	2016	2017
negative	0	0	129.00	0
positive	0	0	31.00	0
$\operatorname{prevalence}(\%)$	NaN	NaN	19.38	NaN

Table 4: Prevalence of Eimeria per year, based on PCR detection. A mouse was considered infected by Eimeria ifone of the 3 markers (COI, 18S or ORF470) gave a sequence, OR if it had a positive count of oocysts in its feces, OR if it was qPCR positive

	2014	2015	2016	2017
negative	0	0	112	0
positive	23	17	48	87
prevalence(%)	100	100	30	100
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Improving Eimeria oocysts detection

22 new samples were detected while diluting by $0.1 \mathrm{mL}$ PBS instead of $1 \mathrm{mL}$ before counting in Neubauer chamber.

Adjusted R-squared = 0.81 represents the amount of variation in y explained by x.

https://www.r-bloggers.com/correlation-and-linear-regression/ (for Lorenzo)

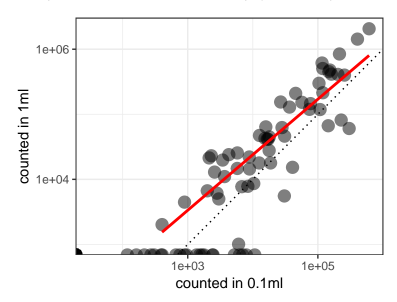


Figure 3: Comparison of OPG depending on dilution level. Red line represents linear relationship between both axis, dotted line represents the function y = x

Comparison oocysts flotation, PCR, qPCR

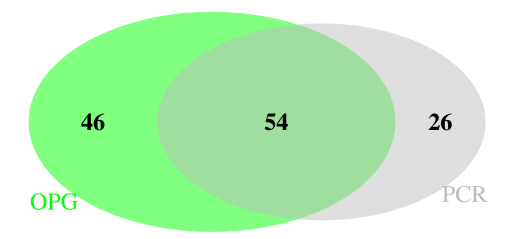


Figure 4: Comparison of detection: PCR vs flotation

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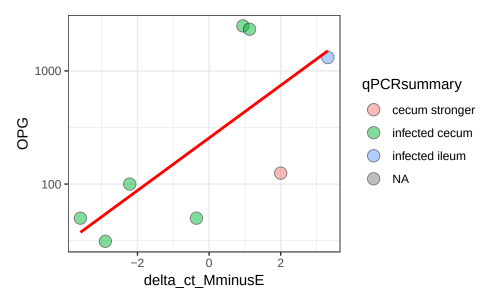


Figure 5: Comparison of positive values of OPG and qPCR for year 2016

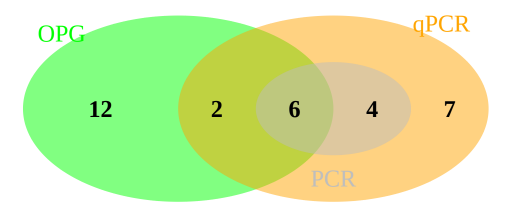


Figure 6: Comparison of detection: PCR vs flotation vs qPCŔ

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Testing hybrid vigor along HMHZ

Oocyst shedding proxy

First approximation:

$geom_smooth()$ using method = 'loess' and formula 'y ~ x'



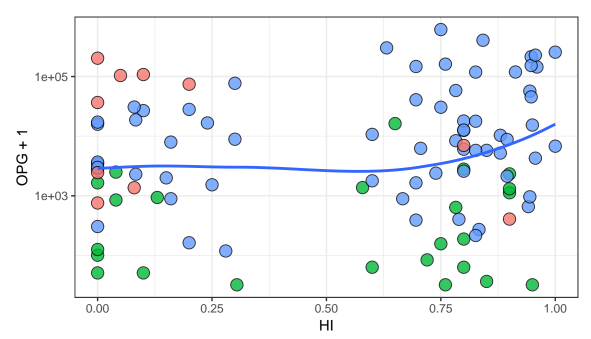


Figure 7: OPG along HI, colored per year. Blue line represent a smooth function (method = loess)

Statistical model (dvp...)

qPCR proxy

tbc

BCI proxy

First approximation:

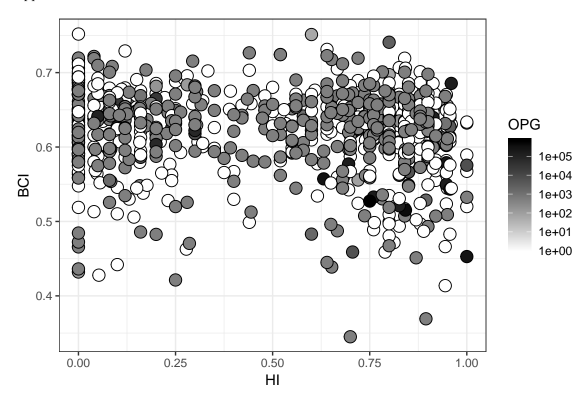


Figure 8: BCI along HI, colored per level of OPG

Bonus part: genotyping of mice case/control

Discussed with Stuart:

- Test distributions 0 or counts. Test all vs only infected ("intensity") distribution. We should be able to fit the distribution of infected on all. Zeros are data. Stochastic move.
- Separation of the zero class. balanced design case/control ~ 400 +/-70infectés SNPchip.
- H0: no differences are observed
- Separate < 0.5 and > 0.5 to see the species effect
- timing: WHEN (for my thesis?)