

Age-dependent susceptiblility to ivermectin and gene expression following an ivermectin blood meal in the malaria vector Anopheles gambiae

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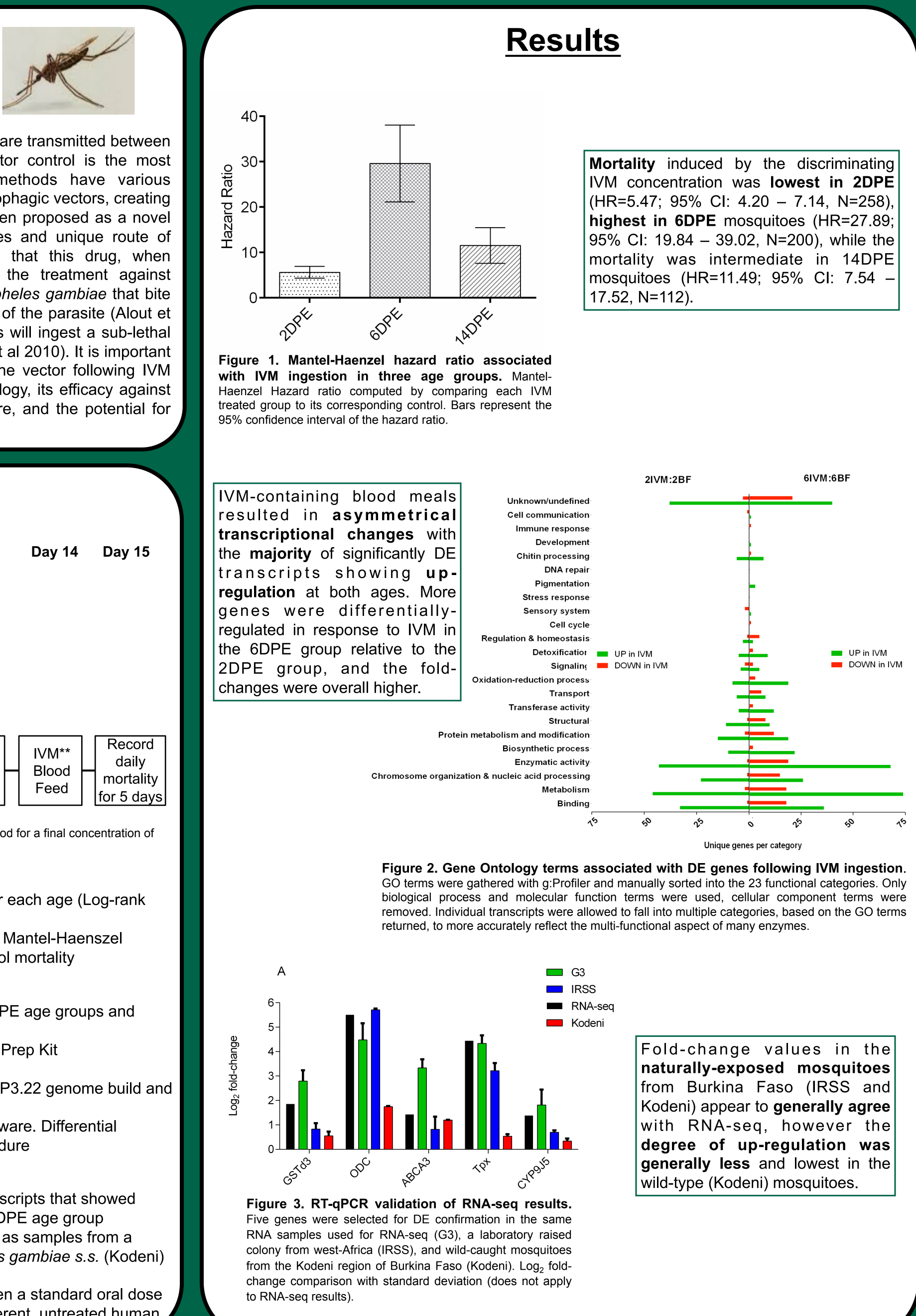
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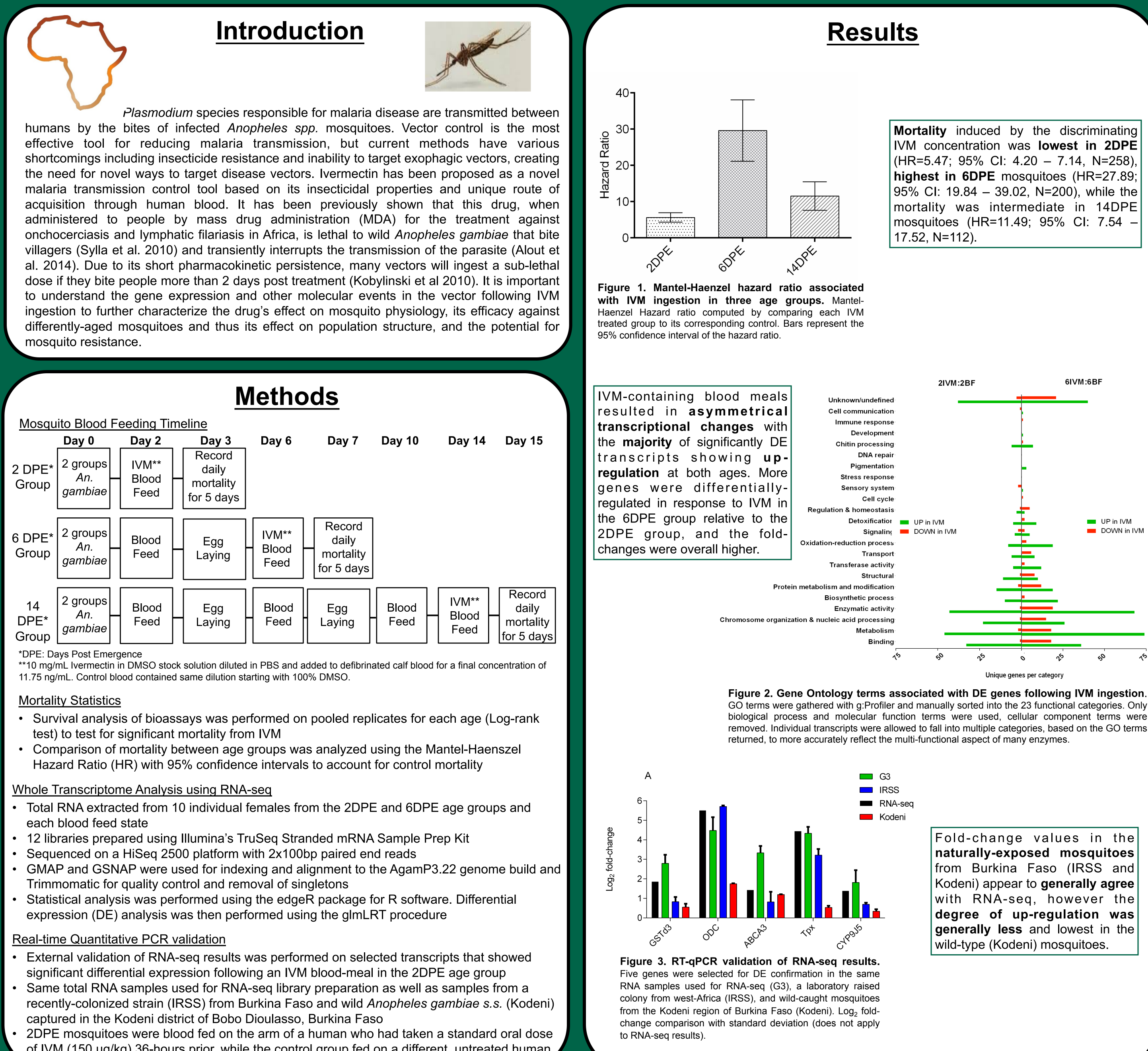
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Age-dependent susceptiblility to ivermectin and gene expression following an ivermectin blood meal in the malaria vector Anopheles gambiae Colorado Jonathan A. Seaman, <u>Haoues Alout</u>, Jacob I. Meyers, Mark D. Stenglein, Roch K. Dabiré, Saul Lozano-Fuentes, Timothy A. Burton, Wojtek S. Kuklinski, William C. Black IV, Brian D. Foy Iniversity Colorado State University, Fort Collins, CO, USA – IRSS, Bobo-Dioulasso, Burkina Faso





- of IVM (150 µg/kg) 36-hours prior, while the control group fed on a different, untreated human.

Genes of interest

Niemann-Pick Type C genes

- AGAP002847 is among the most highly up-regulated genes in response to IVM ingestion at
- 11-fold, respectively)
- evolutionarily distinct from the blood meal-responsive NPC2

Peritrophic matrix associated genes

- had DE genes unique to their own age group.
- feeding but up-regulation of this category after an IVM ingestion
- perhaps contributing to their increased tolerance to IVM toxicity.

Immune response genes

- had DE genes unique to their own age group.

- mosquitoes which have not ingested a prior blood meal.
- explained the age-related susceptibility differences.
- and NPC genes.
- midgut microflora, and innate immune responses.
- groups we have highlighted

Acknowledgements

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Results Cont.

The Niemann-Pick type C-2 (NPC2) family of genes were among the most highly up-regulated transcripts in our experiments in response to blood feeding only and to IVM ingestion

both ages, but is doubled in 6DPE (160-fold) vs. 2DPE (80-fold)".

• AGAP002848 is modestly up-regulated in response to IVM ingestion at both ages (17-fold and

• The IVM-responsive NPC2 genes (AGAP002848 and AGAP002847) appear to be

DE of 45 putative peritrophic matrix-associated genes upon IVM ingestion in both age contrasts suggests that PM gene transcription is responding directly to IVM treatment.

In general the 2DPE contrast showed both an increased number of significantly DE transcripts as well as higher fold-changes than the 6DPE contrast, but interestingly, both age contrasts

Gene ontology analysis revealed down-regulation of chitin processing transcripts upon blood-

The less IVM-susceptible age group (2DPE) shows overall higher DE transcription among PMassociated genes in response to IVM than the more IVM-susceptible age group (6DPE),

The older age contrast showed significant DE among 19 immune-related transcripts and most were modestly up-regulated. Only 4 immune function genes were significantly DE in the 2DPE In general the 2DPE contrast showed both an increased number of significantly DE transcripts as well as higher fold-changes than the 6DPE contrast, but interestingly, both age contrasts

Conclusions

• Our data shown that older An. gambiae mosquitoes that have ingested previous blood meals are more susceptible to IVM compared to young

Following from this difference, **DE analysis showed that IVM in a blood meal** induces significant transcription of more than 100 genes, most of which are up-regulated. There was no single gene or group of genes that clearly

• It is clear that IVM in a blood meal mostly induced transcription of noncanonical genes, including PM-associated genes, immune response genes

The data suggest complex midgut interactions resulting from IVM ingestion that likely involves blood meal digestion physiological responses,

More complex interactions between IVM and the significant DE genes mentioned above are likely as some genes have clear linkages across the