# Supporting Information for:

Elbers JP, Clostio RW and Taylor SS (2016) Population genetic inferences using immune gene SNPs mirror patterns inferred by microsatellites. Molecular Ecology Resources.

**Table S1** Sequencing metrics for *Gopherus polyphemus* samples. Percent UR for percent of total reads that were unique, Percent URA for percent of unique reads that were alignable, Mean coverage for mean number of reads across the target region, Percent 20x for percent of bases in target region with greater than 20x coverage, No. genes for number of genes, and No. exons for number of exons.

	Sample	Total reads	Percent UR	Percent URA	Mean coverage	Percent 20x	No. genes	No. exons
1	AL102	3,212,450	42.5489	98.7624	64.702327	70.9051	592	4,107
2	AL103	$4,\!465,\!410$	41.6559	98.7366	86.236747	74.194	598	$4,\!238$
3	AL106	$3,\!359,\!715$	45.6208	98.8027	71.663432	71.6972	600	$4,\!156$
4	AL108	2,819,070	46.9525	98.4416	61.437226	72.4334	600	$4,\!222$
5	FL846	$3,\!053,\!761$	48.2949	98.7728	68.408722	69.2006	594	$4,\!120$
6	FL855	3,001,641	49.9861	98.8119	70.02287	70.426	587	4,162
7	FL857	$4,\!126,\!014$	48.3302	98.7755	91.599757	73.6741	595	4,209
8	FL880	$2,\!495,\!515$	47.5824	98.5998	55.003413	67.7526	592	4,140
9	GA1044	2,735,000	50.4835	98.8161	64.467663	69.2348	594	$4,\!135$
10	GA1435	$3,\!114,\!664$	48.0062	98.8188	69.54348	69.2231	593	4,088
11	GA1835	$3,\!160,\!564$	47.5032	98.8015	69.893149	70.4834	595	$4,\!135$
12	GA462	1,692,328	50.9997	98.7798	40.600786	61.147	586	3,934
13	LA62	2,490,648	47.5604	98.7996	55.582373	67.2087	592	4,032
14	LA66	$2,\!366,\!268$	48.6917	98.6262	53.254455	65.3586	592	3,992
15	LA77	$3,\!449,\!116$	46.1951	98.7934	74.612173	72.2789	600	$4,\!162$
_16	LA78	1,920,641	55.6212	98.8716	50.102979	63.859	596	3,899

Table S2 All genes with di-allelic, polymorphic SNPs from 16 Gopherus polyphemus samples.

1-phosphatidylinositol 4,5-bisphosphate phosphodiesterase gamma-1-like

16 kDa beta-galactoside-binding lectin-like

25-hydroxyvitamin D-1 alpha hydroxylase, mitochondrial

3-phosphoinositide dependent protein kinase 1

4-hydroxy-2-oxoglutarate aldolase 1

A. superbus venom factor 1-like

acetylcholinesterase-like

active BCR-related

adenosine A2b receptor

adenosine A3 receptor

ADP-ribosylation factor-like 13B

alpha-2-macroglobulin

alpha-2-macroglobulin-like

alpha-2-macroglobulin-like 1

alpha-2-macroglobulin-like protein 1

angiogenin-2-like

ankyrin repeat and death domain containing 1A

annexin A3

antigen-presenting glycoprotein CD1d1-like

apolipoprotein A-IV

apoptosis-associated speck-like protein containing a CARD

aquaporin 4

arrestin, beta 2

ATPase, Cu++ transporting, alpha polypeptide

autophagy related 5

B-cell CLL/lymphoma 10

B-cell CLL/lymphoma 2

B-cell CLL/lymphoma 3

B-cell receptor-associated protein 29

B-cell receptor-associated protein 31

B-cell scaffold protein with ankyrin repeats 1

B and T lymphocyte associated

B lymphoid tyrosine kinase

bactericidal permeability-increasing protein-like

bactericidal/permeability-increasing protein

basic leucine zipper transcription factor, ATF-like

 $beta\hbox{-}2\hbox{-}microglobulin$ 

bone morphogenetic protein 6

bone morphogenetic protein receptor, type IA

breakpoint cluster region

butyrophilin-like protein 9

butyrophilin subfamily 1 member A1-like

butyrophilin subfamily 2 member A1-like

butyrophilin subfamily 3 member A2-like

butyrophilin subfamily 3 member A3-like

C-C chemokine receptor type 5-like

C-C motif chemokine 5-like

C-type lectin domain family 1 member A-like

C-type lectin domain family 2 member B-like

C-type lectin domain family 2 member D-like C-type lectin domain family 4 member D-like C-type lectin domain family 4 member E-like C-type lectin domain family 4 member G-like C-X-C motif chemokine 10-like C3a anaphylatoxin chemotactic receptor-like C4b-binding protein alpha chain-like C5a anaphylatoxin chemotactic receptor 1-like cactin, spliceosome C complex subunit calcium binding and coiled-coil domain 2 calcium channel, voltage-dependent, beta 3 subunit calcium channel, voltage-dependent, beta 4 subunit calcium/calmodulin-dependent protein kinase IV calicin cardiotrophin-like cytokine factor 1 caspase recruitment domain family, member 9 cathepsin G-like Cbl proto-oncogene B, E3 ubiquitin protein ligase CD14 molecule CD180 molecule CD226 molecule CD247 molecule CD274 molecule CD36 molecule (thrombospondin receptor) CD37 molecule CD3e molecule, epsilon (CD3-TCR complex) CD40 molecule, TNF receptor superfamily member 5 CD74 molecule, major histocompatibility complex, class II invariant chain CD79a molecule, immunoglobulin-associated alpha CD79b molecule, immunoglobulin-associated beta CD82 molecule cell adhesion molecule 1 cell adhesion molecule 4 cell division cycle 37 cell division cycle 37-like 1 centromere protein F, 350/400kDa chemokine (C-C motif) receptor 7 chemokine (C-X3-C motif) receptor 1 chromosome unknown open reading frame, human C9orf84 cis-aconitate decarboxylase-like class I histocompatibility antigen, F10 alpha chain-like class II histocompatibility antigen, M alpha chain class II, major histocompatibility complex, transactivator coagulation factor II (thrombin) receptor-like 1 coagulation factor XIII, B polypeptide coenzyme Q10 homolog B (S. cerevisiae) coiled-coil domain containing 142 coiled-coil domain containing 151 coiled-coil domain containing 170

collagen, type III, alpha 1 collectin-46-like collectin sub-family member 12 complement C1r-B subcomponent-like complement C2-like complement C3-like complement component 1, q subcomponent binding protein complement component 1, q subcomponent, A chain complement component 1, r subcomponent complement component 1, s subcomponent complement component 8, alpha polypeptide complement decay-accelerating factor-like complement factor B complement factor H complement factor properdin complement receptor type 2-like CUB and Sushi multiple domains 1 cyclic GMP-AMP synthase-like cytochrome P450 27C1 cytotoxic and regulatory T cell molecule cytotoxic T-lymphocyte-associated protein 4 DEAD (Asp-Glu-Ala-Asp) box polypeptide 58 death-associated protein kinase 1 death-associated protein kinase 2-like death-associated protein kinase 3 death-associated protein kinase 3-like dedicator of cytokinesis 2 dedicator of cytokinesis protein 2-like DEXH (Asp-Glu-X-His) box polypeptide 58 discs, large homolog 1 (Drosophila) dispanin subfamily A member 2b DLA class II histocompatibility antigen, DR-1 beta chain-like DnaJ (Hsp40) homolog, subfamily A, member 3 docking protein 6 dual specificity phosphatase 10 duodenase-1-like dynactin 1 E3 ubiquitin-protein ligase TRIM39-like E3 ubiquitin-protein ligase TRIM39 pseudogene E3 ubiquitin-protein ligase TRIM56-like eomesodermin epiregulin excision repair cross-complementation group 1 exonuclease 1 exosome component 3 extracellular matrix protein 1 family with sequence similarity 105, member A family with sequence similarity 177, member A1 family with sequence similarity 83, member E

Fanconi anemia, complementation group F

Fas cell surface death receptor

Fc fragment of IgE, high affinity I, receptor for; gamma polypeptide

feline Gardner-Rasheed sarcoma viral oncogene homolog

feline sarcoma oncogene

fer (fps/fes related) tyrosine kinase

ficolin-1-like

ficolin-2-like

ficolin (collagen/fibrinogen domain containing) 3

fucokinase

FYN oncogene related to SRC, FGR, YES

G-protein coupled receptor 183-like

G patch domain and KOW motifs

gastrula zinc finger protein XlCGF57.1-like

GATA binding protein 3

glucosaminyl (N-acetyl) transferase 3, mucin type

glutamyl-prolyl-tRNA synthetase

glutathione peroxidase 1

glutathione peroxidase 2 (gastrointestinal)

glycosylphosphatidylinositol specific phospholipase D1

granzyme-like protein 2

GRB2-associated binding protein 2

growth arrest-specific 6

GTP cyclohydrolase 1

GTP cyclohydrolase 1-like

H-2 class II histocompatibility antigen, A-R alpha chain-like

H-2 class II histocompatibility antigen, E-S beta chain-like

H2.0-like homeobox

heat shock 60kDa protein 1 (chaperonin)

hemopexin

heparan sulfate (glucosamine) 3-O-sulfotransferase 4

HLA class II histocompatibility antigen, DP alpha 1 chain-like

HLA class II histocompatibility antigen, DR alpha chain-like

HLA class II histocompatibility antigen, DR beta 5 chain-like

HLA class II histocompatibility antigen, DRB1-15 beta chain-like

immunoresponsive 1 homolog (mouse)

importin 11

indoleamine 2,3-dioxygenase 1

inositol polyphosphate-5-phosphatase, 145kDa

insulin-like growth factor 1 receptor

integrin alpha-L-like

interferon-induced guanylate-binding protein 1

interferon-induced guanylate-binding protein 1-like

interferon-induced protein 44-like

interferon-induced protein with tetratricopeptide repeats 1-like

interferon-induced protein with tetratricopeptide repeats 5

interferon induced transmembrane protein 5

interferon kappa-like

interferon regulatory factor 3

membrane cofactor protein-like

Gene interferon regulatory factor 7 interferon regulatory factor 8 interferon, kappa interleukin-1 receptor-associated kinase 1 interleukin-36 receptor antagonist protein interleukin 1 receptor-like 1 interleukin 12 receptor, beta 1 interleukin 12A interleukin 12B interleukin 18 interleukin 18 receptor 1 interleukin 20 receptor beta interleukin 23 receptor interleukin 27 receptor, alpha interleukin 4 receptor interleukin 6 interleukin 7 receptor ISG15 ubiquitin-like modifier Janus kinase 2 Janus kinase 3 kelch-like family member 24 kelch-like family member 35 kelch-like family member 38 kelch-like family member 6 kelch repeat and BTB (POZ) domain containing 8 keratin, type II cytoskeletal 1-like killer cell lectin-like receptor subfamily B member 1B allele C killer cell lectin-like receptor subfamily F member 1 killer cell lectin-like receptor subfamily G member 1 kynureninase lectin, galactoside-binding, soluble, 2 lectin, galactoside-binding, soluble, 8 leucine rich repeat (in FLII) interacting protein 2 leucine rich repeat containing 70 lymphocyte-specific protein tyrosine kinase lymphocyte cytosolic protein 1 (L-plastin) lymphotoxin alpha lysosomal trafficking regulator Mab-21 domain containing 1 macrophage migration inhibitory factor major histocompatibility complex class I-related gene protein-like mannan-binding lectin serine peptidase 1 mannan-binding lectin serine peptidase 2 mast cell protease 1A-like mast cell protease 3-like mediator of RNA polymerase II transcription subunit 1-like melanotransferrin-like membrane-associated ring finger (C3HC4) 1, E3 ubiquitin protein ligase

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mitochondrial carrier 2 mitogen-activated protein kinase-activated protein kinase 2 mitogen-activated protein kinase kinase 7 mitogen-activated protein kinase kinase kinase 14 mothers against decapentaplegic homolog 6-like mucosa-associated lymphoid tissue lymphoma translocation protein 1-like mucosa associated lymphoid tissue lymphoma translocation gene 1 mutL homolog 1 mutS homolog 6 myelin-oligodendrocyte glycoprotein-like mvosin IF myosin light chain, phosphorylatable, fast skeletal muscle N(alpha)-acetyltransferase 25, NatB auxiliary subunit NACHT, LRR and PYD domains-containing protein 1-like NACHT, LRR and PYD domains-containing protein 12-like natural killer cells antigen CD94-like NCK-associated protein 1-like Nedd4 family interacting protein 1 negative regulator of ubiquitin-like proteins 1 NK2 homeobox 3 NLR family member X1 NLR family, CARD domain containing 5 nuclear factor of activated T-cells, cytoplasmic 2-like nuclear fac. of kappa light polypep. gene enhancer in B-cells 2 (p49/p100) nuclear fac. of kappa light polypep. gene enhancer in B-cells inhibitor, alpha olfactomedin-4-like olfactomedin-like olfactomedin 4 oncoprotein induced transcript 3 opioid receptor, kappa 1 OTU deubiquitinase 7B OTU deubiquitinase with linear linkage specificity ovostatin-like paired amphipathic helix protein Sin3a-like PAX interacting (with transcription-activation domain) protein 1 peptidase domain containing associated with muscle regeneration 1 peptidoglycan recognition protein 2 peptidoglycan recognition protein 3-like peroxisome proliferator-activated receptor gamma phosphatidylinositol 4-kinase type 2 alpha phosphodiesterase 4B, cAMP-specific phosphodiesterase 4C, cAMP-specific phosphodiesterase 4D, cAMP-specific phosphoinositide-3-kinase adaptor protein 1 phospholipase A2, group IB (pancreas) phospholipase A2, minor isoenzyme-like phospholipase C, gamma 1 phospholipase D2-like phospholipid scramblase 2

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phospholipid scramblase family, member 5
pleckstrin hom. domain contain., family A (phosphoinositide bind. spec.) member 1
poliovirus receptor-related 2 (herpesvirus entry mediator B)
poly (ADP-ribose) polymerase family, member 9
poly [ADP-ribose] polymerase 9-like
polymerase (RNA) III (DNA directed) polypeptide C (62kD)
polymerase (RNA) III (DNA directed) polypeptide D, 44kDa
polymerase (RNA) III (DNA directed) polypeptide F, 39 kDa
polymerase (RNA) III (DNA directed) polypeptide G (32kD)
polymerase (RNA) III (DNA directed) polypeptide G (32kD)-like
potassium channel tetramerization domain containing 7
POU class 2 homeobox 2
presenilin 2
programmed cell death 1 ligand 2-like
proteasome (prosome, macropain) subunit, beta type, 4
protein kinase C, beta
protein kinase C, delta
protein kinase C, epsilon
protein kinase C, theta
protein kinase C, zeta
protein kinase D2
protein phosphatase 2, regulatory subunit B' ' gamma
protein phosphatase 3, catalytic subunit, beta isozyme
protein tyrosine phosphatase, non-receptor type 2
protein tyrosine phosphatase, non-receptor type 22 (lymphoid)
protein tyrosine phosphatase, non-receptor type 6
protein tyrosine phosphatase, receptor type, C
protein tyrosine phosphatase, receptor type, N polypeptide 2
purine nucleoside phosphorylase
purinergic receptor P2X, ligand-gated ion channel, 7
pyroglutamyl-peptidase I-like
RAB guanine nucleotide exchange factor (GEF) 1
RAB17, member RAS oncogene family
RAB27A, member RAS oncogene family
RanBP-type and C3HC4-type zinc finger containing 1
rano class II histocompatibility antigen, A beta chain-like
RAR-related orphan receptor C
RAS guanvl releasing protein 1 (calcium and DAG-regulated)
receptor-interacting serine-threonine kinase 2
receptor-interacting serine/threonine-protein kinase 2-like
regulator of cell cycle
retinoic acid receptor, alpha
Rho guanine nucleotide exchange factor (GEF) 37
RIB43A domain with coiled-coils 1
ribonuclease-like
ribosomal protein L13a
ribosomal protein S6 kinase, 90kDa, polypeptide 3
ribosomal protein S6 kinase, 90kDa, polypeptide 6
ring finger and CCCH-type domains 1
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ring finger and CCCH-type domains 2
ring finger protein 135
ring finger protein 19B
ring finger protein 31
S100 calcium binding protein A14
SAM and SH3 domain-containing protein 1
SAM and SH3 domain containing 3
SAM domain and HD domain 1
SAM domain, SH3 domain and nuclear localization signals 1
Sec61 alpha 1 subunit (S. cerevisiae)
sema domain, Ig, TM and short cytoplasmic domain, (semaphorin) 4A
sema domain, Ig, TM and short cytoplasmic domain, (semaphorin) 4B
semaphorin-7A-like
semaphorin 7A, GPI membrane anchor (John Milton Hagen blood group)
serine/threonine kinase 11
serpin peptidase inhibitor, clade G (C1 inhibitor), member 1
SH2 domain containing 1A
SH2 domain containing 1B
SH2B adaptor protein 2
sialidase 2 (cytosolic sialidase)
signal transducer and activator of transcription 6, interleukin-4 induced
SIN3 transcription regulator family member A
sirtuin 1
SMAD family member 6
solute carrier family 26 (anion exchanger), member 6
solute carrier family 26 (anion exchanger), member 8
solute carrier family 26 (anion exchanger), member 9
solute carrier family 26, member 10
solute carrier family 30 (zinc transporter), member 8
spinster homolog 1 (Drosophila)
spinster homolog 2 (Drosophila)
spinster homolog 3 (Drosophila)
spleen tyrosine kinase
src kinase associated phosphoprotein 1
src kinase associated phosphoprotein 2
steroidogenic acute regulatory protein
steroidogenic acute regulatory protein, mitochondrial-like
stomatin (EPB72)-like 2
strawberry notch homolog 1 (Drosophila)
strawberry notch homolog 2 (Drosophila)
suppressor of cytokine signaling 5
suppressor of cytokine signaling 5-like
suppressor of Ty 6 homolog (S. cerevisiae)
suppressor of variegation 3-9 homolog 1 (Drosophila)
SWAP switching B-cell complex 70kDa subunit
synaptotagmin binding, cytoplasmic RNA interacting protein
syntaxin 11
syntaxin 19
syntaxin binding protein 2
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Gene
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syntaxin binding protein 3 T-box 21 TANK-binding kinase 1 tec protein tyrosine kinase tetraspanin 19 thaicobrin-like thrombospondin 1 thymocyte selection associated thymocyte selection associated family member 2 TLR4 interactor with leucine-rich repeats TNF receptor-associated factor 1-like TNF receptor-associated factor 2 TNF receptor-associated factor 3 TNF receptor-associated factor 6, E3 ubiquitin protein ligase TNFAIP3 interacting protein 2 TNFAIP3 interacting protein 3 toll-interleukin 1 receptor (TIR) domain containing adaptor protein toll-like receptor 13 toll-like receptor 2 toll-like receptor 7 toll-like receptor 8 toll-like receptor adaptor molecule 1 toll-like receptor adaptor molecule 2 transferrintransforming growth factor, beta 2 transforming growth factor, beta 3 transforming growth factor, beta receptor III transforming growth factor, beta receptor III-like transient receptor potential cation channel subfamily M member 4-like transient receptor potential cation channel, subfamily M, member 4 translation machinery associated 16 homolog (S. cerevisiae) transmembrane protein 125 transmembrane protein 167A tripartite motif-containing protein 10-like tripartite motif-containing protein 7-like tripartite motif containing 56 tumor necrosis factor (ligand) superfamily, member 13b tumor necrosis factor receptor superfamily member 14-like tumor necrosis factor receptor superfamily member 5-like tumor necrosis factor receptor superfamily, member 14 tumor necrosis factor, alpha-induced protein 3 TXK tyrosine kinase tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta UBA domain containing 2 unc-13 homolog D (C. elegans) unc-93 homolog B1 (C. elegans) uncharacterized LOC101938270 uncharacterized LOC101938480 uncharacterized LOC101940718

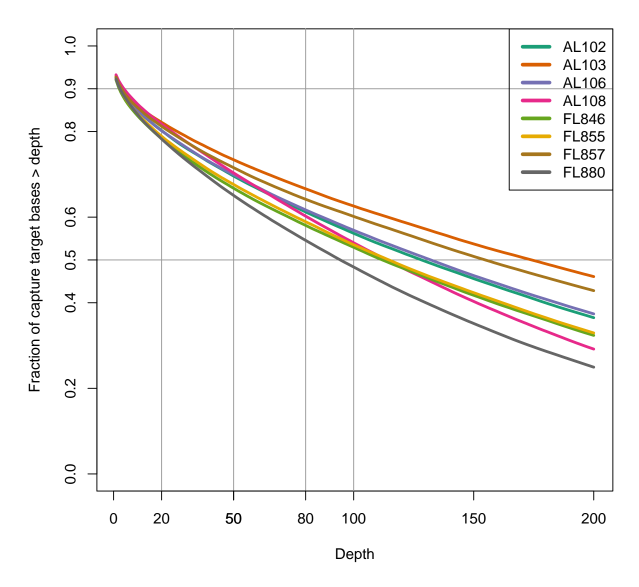
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Table S2 – Continued from previous page

	Table 32 – Continued from previous page
_	Gene
	uncharacterized LOC101941368
	uncharacterized LOC101944094
	uncharacterized LOC101945191
	uncharacterized LOC101945921
	uncharacterized LOC101945971
	uncharacterized LOC101948866
	uncharacterized LOC101948974
	uncharacterized LOC101949200
	uncharacterized LOC101949947
	uncharacterized LOC101950806
	uncharacterized LOC101950941
	uncharacterized LOC101950982
	uncharacterized LOC101951626
	uncharacterized LOC103305939
	uncharacterized LOC103305969
	uncharacterized LOC103305996
	uncharacterized LOC103306364
	uncharacterized LOC103306443
	uncharacterized LOC103306956
	uncharacterized LOC103307015
	uncharacterized protein-like
	v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene homolog
	v-rel avian reticuloendotheliosis viral oncogene homolog A
	v-rel avian reticuloendotheliosis viral oncogene homolog B
	v-yes-1 Yamaguchi sarcoma viral related oncogene homolog
	vav 1 guanine nucleotide exchange factor
	vav 3 guanine nucleotide exchange factor
	VCP-interacting membrane protein
	veficolin-1-like
	venom factor-like
	wingless-type MMTV integration site family, member 5A
	Wiskott-Aldrich syndrome
	X-ray repair complementing defective repair in Chinese hamster cells 4
	Z-DNA binding protein 1
	zeta-chain (TCR) associated protein kinase 70kDa
	zinc-binding protein A33-like
	zinc finger and BTB domain containing 41
	zinc finger protein 239-like
	zinc finger protein 271-like
	zinc finger protein 418-like
	zinc finger protein 436-like
	zinc finger protein 501-like
	zinc finger protein 551-like
	zinc finger protein 572-like
	zinc finger protein 850-like
	zinc finger protein 883-like
	zinc finger protein RFP-like
	zinc finger, SWIM-type containing 7
	zona pellucida sperm-binding protein 3-like
-	

Table S2 - Continued from previous page

zona pellucida sperm-binding protein 4-like



**Fig. S1** Coverage plots for first eight *Gopherus polyphemus* samples showing number of sequencing reads at or above specified proportions. A value at 100 Depth and 0.5 fraction means 50 percent of bases were at or above 100X coverage.

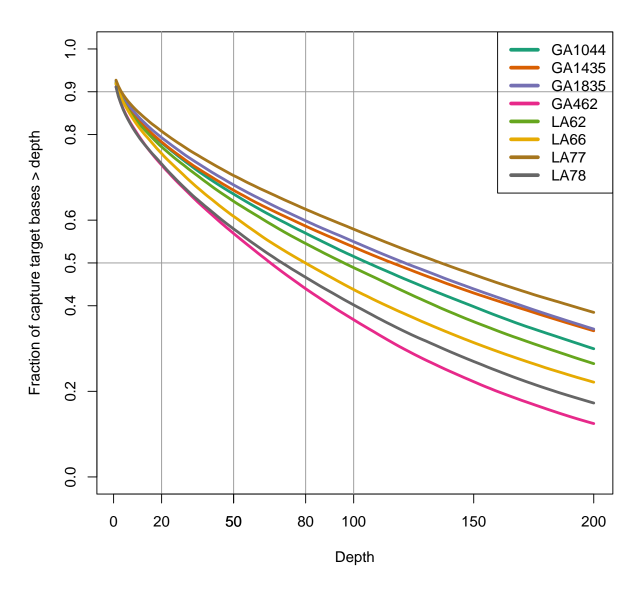


Fig. S2 Coverage plots for last eight *Gopherus polyphemus* samples showing number of sequencing reads at or above specified proportions.

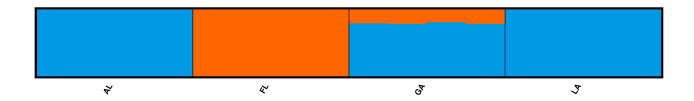


Fig. S3 STRUCTURE plot for 16 Gopherus polyphemus sequenced at 17,901 immune gene SNPs with optimum number of clusters K=2 determined by STRUCTURE HARVESTER.

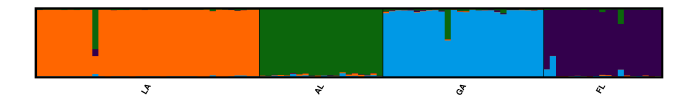


Fig. S4 STRUCTURE plot for the full microsatellite dataset (101 Gopherus polyphemus genotyed at 10 microsatellite loci) with optimum number of clusters K=4 determined by STRUCTURE HARVESTER.



Fig. S5 STRUCTURE plot for the partial microsatellite dataset (16 Gopherus polyphemus genotyed at 10 microsatellite loci) with optimum number of clusters K=3 determined by STRUCTURE HARVESTER.

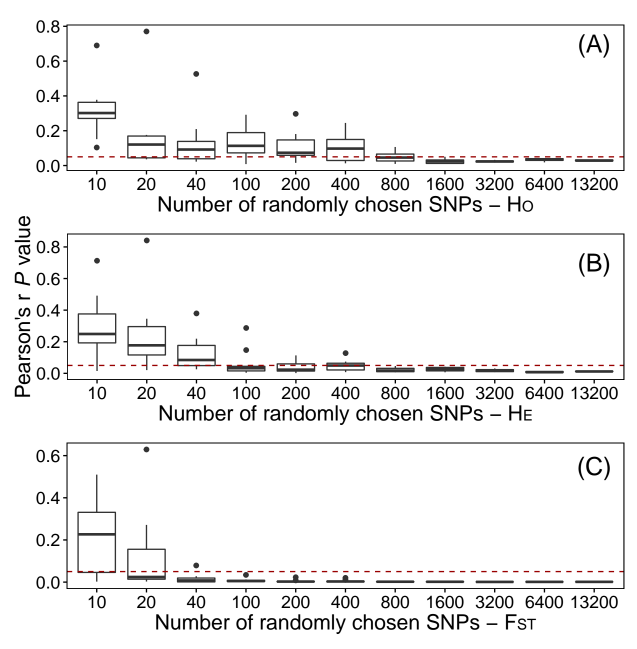


Fig. S6 Subsampling analysis showing how many randomly sampled SNP loci out of the total of 17,901 are needed in comparison to the full microsatellite dataset (101 *Gopherus polyphemus* genotyed at 10 microsatellite loci) for Pearon's r correlation coefficient to be significant at 0.05 level (dotted line) for (A) observed heterozygosity; (B) expected heterozygosity; and (C) FST. There were 10 simulations for each size class of SNPs. Ho for observed heterozygosity, HE for expected heterozygosity.

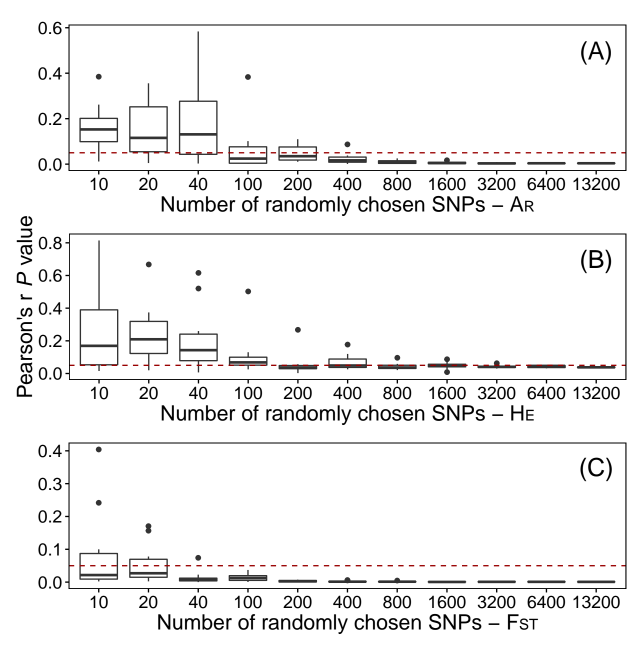


Fig. S7 Subsampling analysis showing how many randomly sampled SNP loci out of the total of 17,901 are needed in comparison to the partial microsatellite dataset (16 Gopherus polyphemus genotyed at 10 microsatellite loci) for Pearon's r correlation coefficient to be significant at 0.05 level (dotted line) for (A) allelic richness; (B) expected heterozygosity; and (C) FST. There were 10 simulations for each size class of SNPs. AR for allelic richness, HE for expected heterozygosity.

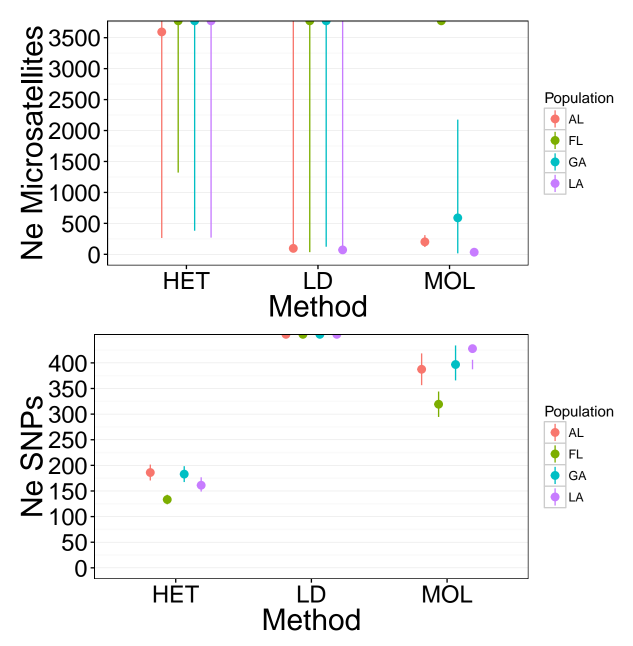


Fig. S8 Effective population sizes per generation (Ne) along with 95 % confidence intervals for Gopherus polyphemus samples estimated with the program NeEstimator using (A) the full microsatellite dataset (101 G. polyphemus genotyed at 10 microsatellite loci) or (B) the SNP dataset (16 G. polyphemus sequenced at 17,901 immune gene SNPs). Dots that are on the top of the graph represent Ne estimates of infinity, and lines that extend to the top of the graph represent upper 95 % confidence limits of infinity. LD for linkage disequilibrium method of Waples & Do (2008), HET for heterozygote-excess method of Zhdanova & Pudovkin (2008), and MOL for the molecular coancestry method of Nomura (2008). Note that the HET and MOL methods estimate the effective number of breeders per year (Nb), which were converted to Ne by multiplying Nb by the generation time of 31 years for G. polyphemus (Enge et al. 2006).