Adaptive genetic variation, stress and glucose regulation

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SUMMARY

Elevated glucose levels in the presence of insulin are indicative of type 2 diabetes and the more inclusive metabolic syndrome. Alleles conferring susceptibility to these and other common conditions may be adaptations to past environments. It is possible that other mammals exhibiting environmental diversity harbor similar variants; therefore, we assessed glucose regulation in two species of deer mice (Peromyscus), a diverse endemic North American group. The prairie deer mouse, P. maniculatus bairdii (BW), and the Oldfield mouse, P. polionotus subgriseus (PO) differ in sexual dimorphism, behavior and habitat. PO animals exhibit better regulatory ability than BW animals, particularly among males, although both species display equivalent insulin levels/responses and non-fasted glucose levels. Hybrid males exhibit a PO glucose challenge response and subsequent dimorphism, behavior and habitat. PO animals exhibit better regulatory ability than BW animals, particularly among males, although both species are known to affect blood pressure and adipose deposition. PO males have GC levels that are twice those of BW males, indicating the presence of alleles that attenuate the GC response. We hypothesize that the interspecific physiological and behavioral differences are interrelated and that similar human variants exist.

INTRODUCTION

The complexity of many prevalent human diseases has led to the suggestion that moderate risk alleles at multiple loci may be a regular occurrence (Smith and Luís, 2002). Termed the common disease/common variant (CD/CV) hypothesis, emerging data has generally supported this model (Peng and Kimmel, 2007; Spielman et al., 2007). Explanations for the CD/CV hypothesis (i.e. high frequency of disease susceptibility alleles) include a lack of negative selection (e.g. diseases which arise later than the average lifespan of pre-civilization humans) or adaptations to past environments. For example, multiple adaptive scenarios have been proposed to explain the rapid increase in type 2 diabetes mellitus (T2DM) and the more inclusive ‘metabolic syndrome’ (Freeman and Cox, 2006; Diamond, 2003; Speakman, 2007) – the latter also includes cardiovascular disease (CVD), stroke and nonalcoholic fatty liver disease. Several lines of evidence suggest the interrelated nature of these diseases (Zimmet et al., 2001). Although obesity-induced metabolic alterations are a cause of these diseases, symptoms may precede obesity and ~20% of T2DM cases are not accompanied by obesity (www.niddk.gov). Given that there are multiple pathways involved in T2DM/metabolic syndrome, it seems likely that multiple relevant environmental factors have changed during the development of human civilization. Stress pathways, for example, are known to affect blood pressure and adipose deposition (Widgren et al., 1992), and to influence CVD (Alevizaki et al., 2007).

If a significant proportion of the human genetic variation involved in disease susceptibility was adaptive, then similar variation may exist in other mammalian species that are adapted to a variety of environments. Such models could be used to identify and study interactions among alleles conferring disease susceptibility in humans. Commonly used laboratory rodents and domesticated animals are not ideal for such studies because they do not represent naturally occurring populations (owing to both allelic combinations and homozygosity) (Beck et al., 2000; Smale et al., 2005).

Deer mice (Peromyscus) are the most common native North American mammals and inhabit nearly every terrestrial habitat on the continent (Dewey and Dawson, 2001). A study of five Peromyscus species suggested a correlation between metabolic rate and local caloric availability (environmental productivity) (Mueller and Diamond, 2001). For example, the California mouse (Peromyscus californicus) is found in chaparral, a relatively unproductive habitat, and has a tendency to develop fatty liver disease, pancreatic pathologies and other T2DM symptoms when given a high-fat diet (Krugner-Higby et al., 2000). These symptoms occur without significant obesity or increase in food intake. It may be relevant that P. californicus exhibits among the strictest monogamy of any mammal species (Ribble, 1991). Monogamy is a rare adaptation with pleiotropic consequences (Clutton-Brock, 1989). Like many of the pathways involved in T2DM/metabolic syndrome, monogamy/pair bonding is at least partly regulated by the hypothalamic-pituitary-adrenal (HPA) axis (DeVries et al., 1995a; Good et al., 2005; Kramer et al., 2005; Taymans et al., 1997). Further, the social interactions required for the pair-bonding observed in monogamy induce stress (Chrousos and Kino, 2007; DeVries et al., 1995b).

Another Peromyscus species, P. polionotus (‘Oldfield mouse’), offers the potential for genetic analysis of monogamy-associated pathways and assessment of their influence on disease phenotypes. P. polionotus has also been documented as monogamous in the wild (Foltz, 1981) and is part of the larger P. maniculatus species.
complex – the best studied of this group is the prairie deer mouse, *P. bairdii*. Captive stocks derived from single wild populations are available for both *P. polionotus* and *P. maniculatus* (Fig. 1). These stocks retain ancestral behavioral differences (e.g. the degree of paternal care of offspring) (Dewey and Dawson, 2001; Kramer et al., 2005; Margulis, 1998).

Both captive and wild-trapped *P. polionotus* (captive stock=PO) animals display a calmer demeanor than their *P. maniculatus* (captive stock=BW) counterparts (Martin et al., 2007). Published data suggest that basal metabolic rates are approximately equal in both the more active, wider ranging *P. maniculatus* and the calmer, monogamous *P. polionotus* (Layne and Kirkland, 1989). Despite being mildly growth-retarded, the offspring of BW females crossed with PO males are healthy and fertile (Dawson et al., 1993). Therefore, these hybrids allow genetic analysis through backcrosses and/or intercrosses. In this system, analysis is facilitated by a nascent genetic map and immanent genome sequencing (Ramsdell et al., 2008).

Because *P. polionotus* and *P. maniculatus* differ in terms of their environment, and both social and non-social behaviors, we hypothesized that the two groups might differ in their susceptibility to T2DM/metabolic syndrome. First, we asked whether the two species were equally adept at glucose homeostasis, a primary diagnostic tool for these diseases.

**RESULTS**

**Species and sex differences in fasting glucose regulatory ability**

We performed glucose tolerance tests (GTTs) on at least 13 individuals of each sex from both the PO and BW stocks. Animals were fasted for 18 hours and then subjected to glucose administration (1.5 mg/g bodyweight) via intraperitoneal injection. We measured blood glucose concentration immediately prior to the glucose challenge (time 0) and every 30 minutes thereafter up until 180 minutes post-administration. All tests were carried out at the same time of day (2-4 p.m.) to avoid confounding effects of circadian rhythms. BW females exhibited significantly higher glucose concentrations (as assessed using the Student’s t-test) than PO females, both at time 0 and throughout most of the measuring period, and there was little overlap between the two groups (Fig. 2A; supplementary material Fig. S1A). Despite these differences, females of both species showed a return to pre-injection blood glucose levels by 90 minutes.

A much larger difference was observed between males of the two species (Fig. 2A; supplementary material Fig. S1B). Initial blood glucose levels for BW males were also well above those of their PO counterparts. As expected, male blood glucose levels rose during the first 30 minutes post-injection and then declined over the next two 30-minute intervals. However, BW males showed only a minimal decline, whereas PO blood glucose concentrations returned to near baseline levels. As shown in Fig. 2A, male BW glucose values remained at almost double the pre-GTT levels, whereas the glucose levels of PO males remained static from 90 minutes after glucose administration. Therefore, the GTT data indicate that Peromyscus glucose regulation exhibits both sex- and species-specific differences.

In comparison with BW males, the male PO GTT response was more similar to the conspecific female response in both shape and absolute values (supplementary material Table S1; supplementary material Fig. S1C,D). For example, values for PO males differed from those of PO females by an average of only 39.6 mg/dL across all time points, whereas the BW sexes differed by an average of 71.5 mg/dL. Together, the pre-GTT data from both sexes suggest either a differential response to fasting or species-specific differences in baseline blood glucose concentrations.

**Genetics of species differences in glucose regulation**

The fertile offspring produced by mating BW females with PO males offer the opportunity for genetic analyses in this system. We performed GTTs on both male and female F1 hybrids to assess the genetics of the species differences in glucose regulation. Female hybrids were not significantly different from either parental strain, suggesting no species dominance (Fig. 2B). In contrast, hybrid males displayed a response that was indistinguishable from their PO fathers (Fig. 2C).

**Species equivalence in male insulin levels and insulin response**

We focused on the differences in male glucose regulation owing to their greater magnitude. We postulated that the differences in male Peromyscus glucose regulation might be the result of different concentrations of circulating insulin. To test this hypothesis, we performed insulin ELISA assays on fasted males before and after glucose administration. These data revealed that the insulin levels in the two strains were equivalent at both time points (Fig. 3A). Computing insulin sensitivity indices based on fasted insulin and glucose values via the QUICKI formula (Katz et al., 2000) resulted in nearly identical values (BW=0.33, PO=0.34).

We also considered that a differential insulin response might underlie the interspecific male variation. We therefore performed insulin tolerance tests (ITTs). The ITT procedure is similar to the GTT, except that insulin is administered rather than glucose. Therefore, blood glucose concentration is expected to initially drop and then return to the pre-test range. Insulin resistance, commonly
associated with T2DM, is marked by a failure of blood glucose levels to fall after the administration of exogenous insulin. We assayed blood glucose concentrations at 15-minute intervals. The two species again displayed significantly different mean glucose concentrations after fasting (Fig. 3B). Females of the two species differed significantly only at time 0 (supplementary material Fig. S2A) and values for PO and BW males exhibited the same pattern. F1 males differed from both PO and BW at 15 minutes post-insulin administration ($P<0.05$) but were otherwise statistically equivalent (supplementary material Fig. S2B). The Peromyscus ITT curves are reminiscent of mild insulin resistance because the animals did not experience dramatic decreases in blood glucose levels. However, it has previously been shown that Peromyscus and some other rodents are not induced to feed by addition of physiological levels of insulin, or by glucose anti-metabolites (Rowland et al., 1985). These data suggest that neither differential insulin concentrations nor insulin sensitivity are the causes of the species differences in glucose homeostasis.

### GTT and ITT responses of Y chromosome consomic animals

The apparent paternal inheritance of glucose homeostasis ability suggested several possibilities: (1) dominance of PO alleles at one or more loci; (2) one or more paternally expressed imprinted loci; or (3) Y chromosome-linked sequences underlie the species differences in glucose regulation. We were able to test the latter hypothesis by breeding consomic animals that had a PO Y chromosome on an otherwise BW genetic background (BW YPO).

The BW YPO animals had significantly different GTT values from BW males at every time point (Fig. 4A). Although more similar to PO males, the consomic animals also differed from them at several time points. The fasted BW YPO animals had lower glucose values than PO males ($P=0.03$), but at 30 minutes post-glucose administration they displayed values that were intermediate between PO and BW (with mean values significantly different from both). From 60 to 120 minutes the consomic pattern was not...
distinguishable from PO values; however, at 150 and 180 minutes post-glucose administration the consomic animals again displayed significantly lower values than PO males. Both PO and BW YPO values at 180 minutes were indistinguishable from their starting blood glucose levels, unlike BW males, whose values were higher than the fasting values.

We also performed ITTs on the consomic animals, which exhibited values within the ranges of both parental strain males (Fig. 4B). Again, the consomic animals displayed starting values that were equivalent to PO males, but diverged to lower values thereafter. These data suggest there may be interactions between the PO Y chromosome and BW autosomal alleles that result in increased insulin sensitivity and/or levels. Testing the effects of the reciprocal combination, a BW Y chromosome on a PO genetic background, is problematic. The difficulty in producing this genetic combination lies in the non-viability of offspring produced by PO females mated to BW males (Duselis and Vrana, 2007).

**Fasting and procedural stress contribute to differences in glucose regulatory ability**

To assess whether the relatively long duration of fasting (18 hours) was responsible for the differences in male response, we conducted GTTs after a 6-hour fast. Differences in the response were clearly present at this time, but were attenuated relative to the 18-hour fast data (supplementary material Fig. S3). However, pre-administration blood glucose levels for both PO and BW males were equivalent to the conspecific levels observed after 18-hour fasts. We also assessed glucose concentration in animals that were fed ad libitum. Without induced fasting or exogenous glucocorticoid administration, males of the two species did not have significantly different mean blood glucose values (~80-100 mg/dL) (supplementary material Table S1). That is, both non-fasted PO and BW males had approximately the same blood glucose concentrations as seen in fasted PO males. Indeed, an entire GTT series performed on non-fasted males indicated no significant differences under these conditions (data not shown). These data suggested that fasting-induced stress might be responsible for a substantial portion of the interspecific differences in glucose regulation.

We therefore performed a mock GTT, in which animals were fasted and then injected with only saline solution. As no glucose is administered, this experiment is a control for stress arising from fasting and procedures. As expected, neither PO nor BW males showed a sharp rise in blood glucose levels during the first 30-minute interval after the saline injection (Fig. 5A). However, BW blood glucose levels rose slowly for approximately 90 minutes thereafter and remained at elevated levels. In contrast, the blood glucose levels did not rise in PO mice. We also tested *P. polionotus* males derived from a distinct and more recently captured population (Santa Rosa Island FL; LS stock) (Lacy and Ballou, 1998). The LS males exhibited the same stability in glucose levels as their PO conspecifics (supplementary material Fig. S2B and data not shown).

We performed the mock GTT on the consomic males (BW YPO), and on both PO and BW females. Plotting this saline response as a percentage of the starting glucose value for each group generated a relatively flat line (transient elevations of ≤140%) for consomic and PO males as well as females of both species. In contrast, BW males reach and remain at ~250% of baseline glucose levels (Fig. 5B). We reanalyzed the GTT data to assess the extent to which the differential stress response underlies the differences in blood glucose concentration. We normalized these data using the values obtained with saline injection only, and compared them to consomic data treated in the same fashion. The normalized patterns are nearly identical, suggesting that the Y chromosome-mediated differences in glucose regulation are largely owing to a differential stress response (supplementary material Fig. S4). These data strongly suggest that there are species-specific aspects of glucose regulation that are influenced by a differential response to both fasting and procedural stress (i.e. handling, transport, injection).

**Evidence that differential glucocorticoid sensitivity results in the species-specific glucose homeostasis**

The effects of stress are mediated by the HPA axis via glucocorticoids (GCs; e.g. cortisol in primates, corticosterone in rodents). These hormones are known to have major effects on glucose metabolism, including stimulation of glycogen synthesis, gluconeogenesis and inhibition of insulin secretion (Lambillotte et al., 1997; Tomlinson and Stewart, 2007). Although elevated GC levels are associated with diabetic or pre-diabetic phenotypes, it is
not clear whether this is always the case. For example, another monogamous mammal, the vole *Microtus ochrogaster*, has higher GC levels and displays an attenuated GC response relative to a non-monogamous sister species (Taymans et al., 1997). Data from previous studies suggested higher GC levels in the monogamous *P. polionotus* than in *P. maniculatus* (Good et al., 2003; Harper and Austad, 2000). In both cases, the captive PO and BW stocks have been shown to have similar responses/GC levels to their wild counterparts; however, the two species have not been compared in the same study. We therefore assayed plasma corticosterone levels from BW, PO and consomic males both prior to and after the stress regimen of fasting and saline injection. As expected, the stress caused GC levels in all three groups to rise (Fig. 6A). However, prior to stress, PO corticosterone levels were ~1.5 times greater than those in BW animals. After fasting and saline injection, PO levels were closer to double the levels in BW animals. In contrast, the consomic males exhibit GC concentrations of only ~25% of those seen in the BW males. The latter values were surprising in that hybrid males had levels equivalent to BW males (supplementary material Fig. S5).

To functionally test the involvement of GC activity in the glucose regulation disparity, we treated animals with the drug metyrapone. Metyrapone acts as a dose-dependent inhibitor of 11-beta-hydroxylase, an essential enzyme in corticosterone/cortisol production. Treatment of fasted males with metyrapone did not affect blood glucose levels in PO animals but lowered the levels in BW males into the PO range at 60-90 minutes post-injection (Fig. 6B). The delay in this reaction is consistent with the time required for metyrapone to act.

**DISCUSSION**

These data indicate differential abilities between the recently diverged species *P. polionotus* and *P. maniculatus* in post-fasting glucose homeostasis. BW males display diabetic phenotypes such as elevated blood glucose, whereas PO males, and females of both species do not. Consistent with the greater variation in males, the genetic data suggest Y chromosome-linked sequences are primarily responsible for the effects. Both fasting and being subjected to sham procedures without a glucose challenge induced the differences in male glucose regulation, implying that these effects are mediated by an underlying differential stress response. We believe this is the first example of natural variation in stress responses resulting in immediate diabetic phenotypes.
Despite the greater stress sensitivity of BW males with regard to glucose regulation, corticosterone levels are much higher in PO males. However, the effects of metyrapone treatment strongly suggest that this stress pathway involves GC hormones. Together, these data imply that PO males are able to attenuate the effects of GC signaling on glucose homeostasis. As both pre- and post-stress insulin levels are similar between the two groups, we hypothesize that differences in glucagon release and/or gluconeogenesis are responsible for the differences. We therefore predict that variation in a Y-linked locus, whose product regulates these pathways, is primarily responsible for this relative GC resistance. This model does not obviously explain the reduced GC levels observed in the consomic animals; however, other studies have also noted that a better understanding of GC turnover is necessary to understand their role in glucose regulation/insulin sensitivity (Dewbury et al., 2007).

Unidentified Y chromosomal sequences are also involved in stress-induced hyperglycemia and subsequent hyperinsulinemia in C3H/WS/Sjn inbred male mice (Leiter, 1988). Unlike the BW males, these animals develop hyperglycemia only after months of repeated stress and then lose this symptom as insulin levels rise, so do not represent naturally-occurring genotypes (Beck et al., 2000; Ideraabdullah et al., 2004). Unidentified human and rat Y chromosomal sequences have been implicated in the stress response and hypertension, but are largely unexamined with respect to glucose homeostasis (Charchar et al., 2002; Dumas et al., 2000; Strathorn et al., 2005). In addition, several studies have shown an association between cortisol metabolism and insulin resistance that is independent of obesity effects (Reynolds et al., 2001). Thus, this system represents an opportunity to develop a novel CD/CV animal model of T2DM/metabolic syndrome, and further studies of mammals with natural allelic combinations will lead to insights that are not possible with mixed and/or inbred lines (Smale et al., 2005). Despite their relative resistance, male PO corticosterone levels still rise in response to stress. Prior studies using P. polionotus have shown an association between stress and corticosterone levels and increased mortality rates of their first litters (Good et al., 2005). GC signaling is enormously complex; GCs are known to affect many pathways, most of which are subject to tissue-specific regulation (Buckingham, 2006; Chrousos and Kino, 2007). Even within the central nervous system, GC signaling appears to differ between the hippocampus and amygdala (Herbert et al., 2006). For example, memory formation and recall have been shown to be differentially affected by increased GC levels (Herbert et al., 2006; Roozendaal et al., 2006). Thus, we predict that increased GC levels in PO animals means that other pathways and organs that are affected by these hormones exhibit greater sensitivity. For example, we predict that the hormone’s effects on glucose homeostasis. This combination results in the calmer demeanor exhibited by the PO males and their ability to maintain normal blood glucose values. Supporting the former hypothesis, addition of exogenous GCs have been shown to aid humans in reducing fear (Soravia et al., 2006).

Despite their relative resistance, male PO corticosterone levels rise in response to stress. Prior studies using P. polionotus have shown an association between stress and corticosterone levels and increased mortality rates of their first litters (Good et al., 2005). GC signaling is enormously complex; GCs are known to affect many pathways, most of which are subject to tissue-specific regulation (Buckingham, 2006; Chrousos and Kino, 2007). Even within the central nervous system, GC signaling appears to differ between the hippocampus and amygdala (Herbert et al., 2006). For example, memory formation and recall have been shown to be differentially affected by increased GC levels (Herbert et al., 2006; Roozendaal et al., 2006). Thus, we predict that increased GC levels in PO animals means that other pathways and organs that are affected by these hormones exhibit greater sensitivity. For example, we predict that the like the similarly monogamous P. californicus, P. polionotus will be more susceptible to the effects of a higher fat diet.

The species differences in the GTT pattern are reminiscent of the observation that monogamy (e.g. as seen in P. polionotus) is associated with a reduction in sex-specific differences. There is a greater similarity in the GTT responses between male and female PO animals than between male and female BW animals. Our hypothesis is that the superior glucose homeostasis ability of the PO males is related to their monogamy.

The relative GC resistance exhibited by PO males strengthens a trend associated with monogamous mammals (Table 1). Within Peromyscus, the monogamous P. eremicus exhibit corticosterone levels that are between 2-4 times higher than those of non-monogamous white-footed mice P. leucopus (Martin et al., 2007). The monogamous prairie vole M. ochrogaster exhibits corticosterone levels that are 8-10 times higher than the recently diverged non-monogamous montane vole M. montanus (Taymans et al., 1997). Increased GC levels and/or GC resistance is also associated with monogamy and/or paternal care of offspring in several primate species (Bales et al., 2006; da Silva Mota et al., 2006).

Monogamy is thought to be an adaptation to those environments that demand greater paternal involvement in offspring care and/or stricter guarding of females against potential rivals (Clutton-Brock, 1989; Marlowe, 2000). Compared with BW males, PO males exhibit both increased paternal care and greater aggressiveness towards unrelated males (Layne and Kirkland, 1989; Margulis, 1998; Martin et al., 2007). We suggest that P. polionotus has adapted to these stressors by increasing corticosterone levels, but attenuating the hormone’s effects on glucose homeostasis. This combination results in the calmer demeanor exhibited by the PO males and their ability to maintain normal blood glucose values. Supporting the former hypothesis, addition of exogenous GCs have been shown to aid humans in reducing fear (Soravia et al., 2006).

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Major efforts in treating T2DM/metabolic syndrome symptoms are focused on the local effects of GC signaling (Tomlinson and Stewart, 2007). Elucidation of the mechanisms employed by P. polionotus to attenuate the effects of GC signaling on glucose homeostasis could significantly aid these efforts. We believe that studying the variation in the P. maniculatus species complex experimental system has a unique potential for understanding the relationship between environment-associated adaptations and mammalian disease susceptibility. In general, we suggest that further studies of mammals with natural allelic combinations will yield insights that are not possible with mixed and/or inbred lines (Smale et al., 2005).

It appears possible that variations in the human stress response may have been shaped by environmental and/or cultural differences. Social mores in much of the world are indicative of a

### Table 1. Same-study glucocorticoid comparisons of monogamous versus promiscuous species/populations

<table>
<thead>
<tr>
<th>Monogamy/paternal care</th>
<th>Promiscuous/lack of care</th>
<th>Relative GC levels</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>P. polionotus</td>
<td>P. maniculatus</td>
<td>Monog ~2× higher</td>
<td>This study</td>
</tr>
<tr>
<td>P. eremicus</td>
<td>P. leucopus</td>
<td>Monog ~2-4× higher</td>
<td>Glaser and DeVries, 2005</td>
</tr>
<tr>
<td>M. ochrogaster</td>
<td>M. montanus</td>
<td>Monog ~5-10× higher</td>
<td>DeVries et al., 1995a; Taymans et al., 1997a</td>
</tr>
<tr>
<td>Leontopithecus rosalia</td>
<td>Leontopithecus rosalia</td>
<td>Monog ~2× higher</td>
<td>Bales et al., 2006</td>
</tr>
<tr>
<td>Callithrix jacchus</td>
<td>Callithrix jacchus</td>
<td>Exp fathers ~2-3× higher</td>
<td>da Silva Mota et al., 2006</td>
</tr>
</tbody>
</table>

Abbreviations: Monog=monogamous; Exp=experienced.
general trend towards social/genetic monogamy in humans (Wyllings et al., 2006). However, cultural variation still exists, and there is evidence that the shift towards monogamy is recent (Dupanloup et al., 2003). Ample evidence indicates roles for GC signaling in modulating sexual/social bonding and other interactions in humans (Curtis et al., 2006; Halpern et al., 2002; Storey et al., 2000; Wyatt et al., 2007). Thus, we predict the existence of GC pathway-related alleles that were selected for in earlier non-monogamous cultures, which now result in increased disease susceptibility.

Detailing the molecular mechanisms by which this specificity is conferred will be a key component of deciphering the relationship between common alleles related to stress/GC influenced pathways and disease. However, we believe that understanding environmental and social influences on these pathways, including distinctions between varieties of social stress, will also be essential.

METHODS

Animals

Origins and husbandry

P. polionotus and P. maniculatus animals were purchased from the Peromyscus Genetic Stock Center (PGSC; http://stkctr.biol.sc.edu/). These animals were caught from single populations in the wild as noted in the text and on the PGSC website. Animals bred at the PGSC or descendents bred at UCI were used in the tests. Animals were housed with food and water ad libitum on a 16:8 hour light:dark cycle. All animals tested were at least 65 days old but not more than 20 months old (both species breed past 2 years of age and live for >4 years). Animals were fed a standard low-fat (4% by weight) rodent chow. We bred BW males and their subsequent male progeny to BW females for eleven generations. The consomic animals tested were from tenth and eleventh generation backcrosses. The sex ratios produced from the offspring of BW females bred to BW YPO males were not significantly different from the PO and BW stocks (supplementary material Fig. S6). We genotyped BW YPO animals at eight unlinked autosomal and X-linked loci, all of which showed the expected BW genotype. The consomic animals were equivalent in size to BW males. Female animals produced by the consomic line were indistinguishable from other BW females in body weight and GTT response.

Assays

Glucose tolerance tests (GTTs)

As noted in the text, adult animals were fasted prior to testing. Animals were weighed and then given an intraperitoneal injection of 1.5 mg glucose per gram of body weight in a saline solution. Blood glucose concentration was measured from a small cut made at the end of the tail. The scab was gently scraped off at subsequent measurements. Approximately 5 μL of blood was then spotted onto commercially available glucose test strips and analyzed using the accompanying glucose meter (Ascensia Elite XL model).

Assays

Insulin tolerance tests (ITTs)

As with the GTT procedure described above, adult animals were fasted prior to testing. The fasting duration was reduced to 8 hours of fasting prior to testing. The fasting duration was reduced to 8 hours.
to reduce mortality during the monitoring period (i.e. owing to hypoglycemic shock). We tested insulin concentrations of both 1 and 0.5 units per kilogram of bodyweight. Insulin was administered and blood glucose measured as described for the GTT procedure.

Stress inhibitor assays
These assays were performed in a similar way to the GTT and ITT procedures described above, except that the indicated drug was administered in a saline solution. The dosages used were: metyrapone, 50 mg/kg bodyweight and propranolol hydrochloride, 10 mg/kg bodyweight.

ELISAs
A corticosterone ELISA kit (Assay Systems, Ann Arbor, MI) was used to measure plasma corticosterone levels as per the instructions of the manufacturer. The insulin ELISA was performed using a kit from Crystal Chem (Downers Grove, IL; Cat. #90060) as per manufacturer’s instructions. All assays were performed on blood from adult animals obtained from the tail. Six or more animals per genotype were analyzed for each assay.

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COMPETING INTERESTS
The authors declare no competing financial interests.

AUTHOR CONTRIBUTIONS
R.C.O., C.D.W., M.J.D. and P.B.V. conceived, designed and performed the experiments. P.B.V. wrote the paper.

SUPPLEMENTARY MATERIAL
Supplementary material for this article is available at http://dmm.biologists.org/content/1/4-5/255/suppl/DC1

REFERENCES


GTT after 6 hour fast

**Blood Glucose (mg/dL)**

- PO male
- BW male
Cortisol (ng/mL)

- BW
- PO
- Consomic
- F1
BW YPO Consomic Line Sex Ratios

Generation

Percent Females

G07  G08  G09  G10  G11  G12  G7-12  PO  BW
### Table S1. Blood glucose values at indicated times (mg/dL)

<table>
<thead>
<tr>
<th>Time</th>
<th>BW female</th>
<th>BW male*</th>
<th>PO female</th>
<th>PO male*</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 (pre-fast)</td>
<td>103.6±20.5</td>
<td>91.9±17.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 (post-fast)</td>
<td></td>
<td>110.3±22.7</td>
<td>121.2±36.3</td>
<td>70.6±18.7</td>
</tr>
<tr>
<td>30</td>
<td>148±50.5</td>
<td>229.5±49.7</td>
<td>82.5±32</td>
<td>136.8±58.8</td>
</tr>
<tr>
<td>60</td>
<td>137.7±68.6</td>
<td>215.4±59.3</td>
<td>75.4±22.9</td>
<td>123.5±41.3</td>
</tr>
<tr>
<td>90</td>
<td>108.5±81.6</td>
<td>207±82.5</td>
<td>72.8±26.5</td>
<td>102.1±22.1</td>
</tr>
<tr>
<td>120</td>
<td>111.8±66.7</td>
<td>210±87.2</td>
<td>67.1±31.2</td>
<td>104.2±30.8</td>
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<td>150</td>
<td>107.1±33.4</td>
<td>172±62.8</td>
<td>57.7±21.2</td>
<td>105.9±30.2</td>
</tr>
<tr>
<td>180</td>
<td>128.8±55.1</td>
<td>197.3±112.9</td>
<td>61.4±16.4</td>
<td>105.1±20.9</td>
</tr>
</tbody>
</table>

*Note that although BW fasting glucose values are always significantly higher than PO values of the same sex, these values do fluctuate over time, which is likely to be dependent on the time of year and/or other external cues.*