Evidence That the Sympathetic Nervous System Elicits Rapid, Coordinated, and Reciprocal Adjustments of Insulin Secretion and Insulin Sensitivity During Cold Exposure

Dynamic adjustment of insulin secretion to compensate for changes of insulin sensitivity that result from alteration of nutritional or metabolic status is a fundamental aspect of glucose homeostasis. To investigate the role of the brain in this coupling process, we used cold exposure as an experimental paradigm because the sympathetic nervous system (SNS) helps to coordinate the major shifts of tissue glucose utilization needed to ensure that increased thermogenic needs are met. We found that glucose-induced insulin secretion declined by 50% in rats housed at 5°C for 28 h, and yet, glucose tolerance did not change, owing to a doubling of insulin sensitivity. These potent effects on insulin secretion and sensitivity were fully reversed by returning animals to room temperature (22°C) for 4 h or by intravenous infusion of the α-adrenergic receptor antagonist phentolamine for only 30 min. By comparison, insulin clearance was not affected by cold exposure or phentolamine infusion. These findings offer direct evidence of a key role for the brain, acting via the SNS, in the rapid, highly coordinated, and reciprocal changes of insulin secretion and insulin sensitivity that preserve glucose homeostasis in the setting of cold exposure.

The capacity to adjust insulin secretion to compensate for changes of systemic insulin sensitivity is an essential but poorly understood aspect of glucose homeostasis (1–3). In insulin-resistant conditions such as obesity (3), pregnancy (4), or adolescence (5), for example, insulin secretion must increase to maintain euglycemia, and the reverse applies to conditions of increased insulin sensitivity (1–3,6). Failure to mount an appropriate pancreatic β-cell response to worsening insulin resistance is a hallmark of the progression from normal glucose tolerance to type 2 diabetes (T2D) (7).

The concept that insulin sensitivity and insulin secretion are reciprocally regulated stems from human studies in which plasma glucose and insulin levels measured during a frequently sampled intravenous glucose tolerance test (FSIGT) were analyzed by the minimal model method. This approach generates reliable estimates of key determinants of glucose tolerance: insulin secretion (typically measured as the acute
The coupling of insulin secretion to insulin sensitivity cannot be explained by changes of plasma glucose levels (8). Although rising nocturnal plasma levels of nonesterified fatty acids (NEFA) were identified as a potential mechanism linking increased insulin secretion to obesity-induced insulin resistance (9), the extent to which this mechanism contributes to the hyperbolic relationship between insulin secretion and insulin sensitivity remains unknown. To investigate the role of the brain in this coupling mechanism, we sought to interrogate its contribution to the adaptive metabolic response to a physiological challenge. To this end, we used cold exposure as a provocative intervention, based on the following considerations: 1) maintenance of core body temperature in cold environments depends on temperature-sensitive hypothalamic neurocircuits that increase sympathetic nervous system (SNS) outflow to thermogenic tissues (10,11); 2) cold exposure increases SNS outflow to the pancreas (12), which likely explains the associated reduction of insulin secretion (via a mechanism involving activation of \( \alpha \)-adrenergic receptors on pancreatic \( \beta \)-cells [13–15]); and 3) glucose tolerance does not change or is improved despite the effect of cold exposure to decrease insulin secretion (16,17).

Although the latter observation can be accounted for by the known effect of cold exposure to increase insulin sensitivity in thermogenic tissues (17–19), the goal of the current work was to determine whether the brain serves to ensure that insulin secretion declines in a manner that preserves glucose homeostasis and averts hypoglycemia. To this end, we measured the effect of cold exposure on determinants of glucose tolerance in rats using the FSIGT/minimal model approach in the presence and absence of an intravenous (i.v.) infusion of the \( \alpha \)-adrenergic receptor (\( \alpha \)-AR) blocker phentolamine. Because the proper interpretation of our data hinges on whether the relationship between insulin secretion and insulin sensitivity in rats comports with the hyperbolic relationship characteristic of humans, we also sought to create a normative rat database with which to interrogate this relationship. This goal was achieved by analyzing pooled data from FSIGT studies conducted in either of two strains of rat (Wistar or Long-Evans) fed a standard chow diet or a high-fat diet (HFD), some of which were included as normal control data in a previous publication (20).

We report that, as expected, the relationship between insulin sensitivity and insulin secretion in rats conforms to the hyperbolic function characteristic of humans. We further demonstrate that the potent effect of cold exposure to the increase of insulin sensitivity is perfectly offset by a proportionate decline of glucose-induced insulin secretion such that glucose tolerance remains unchanged. Moreover, the effects of cold exposure on insulin secretion and insulin sensitivity are rapidly reversed by systemic \( \alpha \)-AR blockade. Together, these findings offer direct evidence of a key role for the brain in the highly coordinated, potent, and reciprocal adjustments of insulin sensitivity and insulin secretion that maintain stable glucose homeostasis during cold exposure.

**RESEARCH DESIGN AND METHODS**

All procedures were performed in accordance with National Institutes of Health Guidelines for the Care and Use of Animals and were approved by the University of Washington Animal Care Committee.

**Experimental Animals**

Adult male Wistar and Long Evans rats (Harlan Laboratories, Indianapolis, IN) were individually housed under specific pathogen-free conditions in a temperature-controlled room with a 12:12 h light/dark cycle and provided ad libitum access to water and standard laboratory chow (PMI Nutrition, St. Louis, MO) or an HFD containing 60% kcal fat (D12492; Research Diets, Inc., New Brunswick, NJ), unless otherwise stated.

**Surgery**

Rats underwent implantation of catheters into the carotid artery and jugular vein and received buprenorphine hydrochloride (Reckitt Colman Pharmaceuticals, Richmond, VA) at the completion of the surgery as previously described (20–22).

**FSIGT and Minimal Model Analysis**

After an overnight fast, multiple blood samples were obtained from unrestrained, conscious animals via an arterial catheter as previously described (20–23). Plasma insulin and blood glucose levels generated from the FSIGTs were analyzed using MinMod software to quantify \( S_C \) and \( S_I \) as previously described (20,23,24). \( \text{AIRG} \), the incremental area under the insulin curve during the FSIGT (AUC\text{insulin}), the BIE, GEZI, and the glucose disappearance rate constant (\( K_G \)) were calculated as previously described (20,23,24). A detailed description of the FSIGT and minimal model analysis is provided in the Supplementary Data.

**Euglycemic-Hyperinsulinemic Clamp**

To independently assess the effect of phentolamine on insulin sensitivity in cold-exposed rats, a euglycemic-hyperinsulinemic clamp was performed as previously described (21,22). A detailed description of the clamp procedure is provided in the Supplementary Data.

**Analysis of Rates of Insulin Secretion and Insulin Clearance**

Insulin clearance was estimated by first calculating the insulin secretion rate using deconvolution of the measured C-peptide concentrations, followed by calculation of the
rate of insulin clearance as AUC(insulin secretion)/AUC (plasma insulin). The parameters for rat C-peptide kinetics were obtained from a kinetic study using 5 nmol/kg injections of rat C-peptide 2 in Sprague-Dawley rats. The C-peptide concentration data after the injection were well described by a one-compartment model with clearance = 29.7 mL/min/kg and volume = 656 mL/kg (25).

A Normative Database for Analysis of Relationships Between Insulin Sensitivity, Insulin Secretion, \(S_D\), and Glucose Tolerance in Rats
To create a normative database with which to analyze relationships between the determinants of glucose tolerance in rats, we assembled data from studies previously conducted in adult male rats in our laboratory that used the FSIGT and minimal model analysis protocol described above that included two different rat strains (Wistar \(n = 54\) or Long-Evans \(n = 25\)), provided ad libitum access to chow \((n = 62)\) or an HFD \((n = 17)\) for 5 days or 3 months before study, vehicle-treated animals from a previous study \((20)\), and animals from the current studies.

A key goal was to determine whether the relationship between insulin sensitivity and insulin secretion conforms to a rectangular hyperbola in rats as it does in humans \((2,3,26,27)\). To test this hypothesis, we used reduced major axis regression (RMA) with bootstrapped estimates of the SEs \((28)\) \((https://www.exetersoftware.com/cat/biomstat/book_biometry.html)\) to fit linear models for the relationship between a measure of insulin secretion (AIR\(_G\)) or an integrated measure of the circulating insulin level during the FSIGT (AUC\(_{insulin}\)), which reflects insulin secretion and its clearance from plasma, and the inverse of insulin sensitivity \((i.e., as a function of [1/S_i])\). For completeness, we also modeled insulin secretion equaling a regression slope times 

\[
\frac{1}{S_i} \times \text{RMA}
\]

among the same temperatures. Experiments were performed using a crossover design, such that each animal was studied at each temperature separated by at least 10 days. To assess the durability of cold-induced changes in determinants of glucose tolerance, separate cohorts of rats \((n = 9/\text{group})\) were housed for 24 h at room temperature or at 5°C, and then housed at room temperature for an additional 4 h before and during an FSIGT.

Role of the SNS on Determinants of Glucose Tolerance in Cold-Exposed Rats
To determine whether \(\alpha-AR\) signaling is required for the effects of cold exposure on AIR\(_G\) and \(S_i\), rats with chronic indwelling catheters were exposed to the cold (5°C) for 28 h without access to food. At 30 min before the FSIGT, a continuous i.v. infusion of the nonselective \(\alpha-AR\) blocker phentolamine at a dose of 8 \(\mu\)g/kg/min (identified in pilot studies as a dose that has minimal effects on mean arterial pressure and heart rate) (Supplementary Fig. 1) or vehicle \((n = 9/\text{group})\) was commenced, with infusion continuing throughout the FSIGT.

Effect of Phentolamine on Determinants of Insulin Action in Cold-Exposed Rats
After implantation of catheters, adult male Wistar rats were exposed to the cold (5°C) for 28 h without access to food, and a euglycemic-hyperinsulinemic clamp was performed \((21,22)\). Although tracer dilution analysis using tritiated glucose was included in the study design, the data are not included owing to technical problems that confound data interpretation. Animals received a continuous i.v. infusion beginning at \(t = -30\) min of phentolamine (8 \(\mu\)g/kg/min) or its vehicle. A primed continuous infusion of regular human insulin (16 mU/kg bolus, followed by 2 mU/kg/min HumulinR; Eli Lilly, Indianapolis, IN) was initiated at \(t = 0\) min. A 50% dextrose solution was infused as required to maintain euglycemia, and serial blood samples were obtained for determination of plasma glucose and insulin levels.

Blood Collection and Assay
Whole blood samples for plasma hormonal measures were collected in appropriately treated tubes \((22)\) and centrifuged, and the plasma was removed and stored at \(-80°C\) for subsequent assay. Plasma glucose was measured using a GM9D glucose direct analyzer (Analox Instruments, Stourbridge, U.K.), plasma insulin (Crystal Chem, Downers Grove, IL), glucagon (Mercodia, Uppsala, Sweden), and corticosterone (Alpco, Salem, NH) by ELISA and NEFAs by colorimetry (Wako Chemicals, Richmond, VA).

Statistical Analysis
Results are expressed as mean ± SEM. Significance was established at \(P < 0.05\) (two-tailed). Statistical analyses of effects of cold exposure or phentolamine on metabolic parameters were performed using Statistica 7.1 software (StatSoft, Inc., Tulsa, OK). A one-way ANOVA with a least significant difference post hoc test was used to compare mean values among multiple groups, a two-sample unpaired Student \(t\) test was used for two-group comparisons, and a paired Student \(t\) test was used for within-group comparisons. RMA regression was performed via constrained nonlinear regression in SPSS 23 software (IBM Corp., Armonk, NY) using sequential quadratic programming and the loss function specified in the IBM technical note available at http://www-01.ibm.com/support/docview.wss?uid=swg21476031. The general linear model in SPSS was used to model the relationships among measures of glucose tolerance and measures of insulin- and noninsulin-mediated determinants of glucose homeostasis.
RESULTS

Relationship Between Insulin Sensitivity and the Insulin Response to Glucose in Rats

Analysis of data compiled from a cohort of 79 normal rats revealed the predicted hyperbolic relationship between insulin sensitivity (SI) and the insulin response to glucose, whether the latter was measured as AIRG or as AUCinsulin (Fig. 1A and B). Although RMA regression identified asymptotic values (y-intercepts) with bootstrap estimated 95% CIs that did not include zero (383, 689 for AIRG; 889, 2673 for AUCinsulin), indicating that the intercepts are statistically significant, inspection of Fig. 1 suggests that the relationship between SI and either measure of the insulin response is fundamentally hyperbolic in nature. We suspect that because AUCinsulin reflects the exposure of body tissues to insulin during the FSIGT more effectively than does AIRG, the relationship between SI and AUCinsulin conforms to a hyperbola more closely than does that between SI and AIRG (Fig. 1A and B). Moreover, the insulin sensitivity of rats fed the HFD was reduced relative to that of chow-fed controls (mean SI: 0.80 ± 0.39 vs. 3.80 ± 2.84 [0⁻⁴ × min⁻¹]/µU/mL; P < 0.0001), whereas insulin secretion was increased (mean AIRG: 2,157 ± 700 vs. 1,003 ± 480 µU/mL × min; P < 0.0001) (Fig. 1C).

To assess the relative contributions of insulin-dependent and insulin-independent mechanisms to glucose disposal during the FSIGT, we performed multiple regression analyses in which the dependent variables were either of two measures of glucose tolerance (K_G or AUCglucose) during the FSIGT. These models included as predictor variables the interaction of the insulin response (measured as AIRG or AUCinsulin) and insulin sensitivity, along with SG (Table 1 and Fig. 1D). As indicated in Table 1, SG makes a substantial contribution to glucose tolerance whether the latter is measured as K_G (models 1 and 2) or AUCglucose (models 3 and 4), as in humans (30). Relatedly, the contribution made by overall insulin action (represented by the DI or by the product of SI × AUCinsulin) to glucose tolerance appears to be larger when AUCglucose is used instead of K_G.

To analyze the relationship between insulin action and glucose tolerance among rats with relatively high or low values of SG, regression lines were fit by ANCOVA for each cohort (Fig. 1D). Although the slopes of the lines fit to each of the two groups did not differ significantly (P = 0.54), for

Figure 1—Relationship between insulin sensitivity and insulin secretion is hyperbolic in nature in rats. Data are from control rats tested at room temperature (22°C; n = 79) depicting the relationships between insulin sensitivity (SI) measured via the minimal model and first-phase insulin secretion measured as AIRG (A); incremental AUCinsulin measured as total AUC during the FSIGT minus basal insulin AUC (B); and AIRG in HFD- and chow-fed rats shown separately (C). D: Simplified depiction of model 4 of Table 1 shows an ANCOVA model for the dependence of glucose tolerance (measured as the net AUCglucose during the FSIGT) on insulin action (measured as SI × net AUCinsulin) in groups categorized by median splits of SG. Solid curves are equations of the form: insulin response = a + b × (1/insulin sensitivity), where the optimal model fit was achieved using RMA regression to account for error in both the x and y variables (RESEARCH DESIGN AND METHODS). Dashed curves are equations of the form: insulin response = c × (1/insulin sensitivity) fit by ordinary least squares optimization using a model that excluded the intercept term in accordance with the concept that insulin secretion × sensitivity ≈ constant for subjects with normal glucose tolerance (the hyperbolic law of glucose homeostasis). For AUCinsulin, the agreement of the optimal RMA fit with the hyperbolic-law curve is excellent, consistent with the fact that insulin acts across the entire glucose tolerance test to promote a return to normoglycemia.
any given value of insulin action, the rats in the upper SC split had significantly better (P < 0.0001) mean overall glucose tolerance than did rats in the lower SC split. These observations suggest that the effect of insulin action on glucose tolerance in rats is influenced by variation in SC, a conclusion consistent with data obtained from three other models (data not shown). Similar outcomes were obtained using the models that replaced SC with GEZI (data not shown), consistent with human data (30). Indeed, GEZI (but not BIE) appears to be a substantial determinant of glucose tolerance in rats, given that the Pearson correlation (r) between Kc and GEZI was 0.56 (P < 0.0001).

Overall, our results indicate that either AIRG or AUCinsulin is a useful index of the insulin response to glucose and that together they support the conclusion that variation in insulin sensitivity across animals is effectively compensated by a proportionate change of insulin secretion.

Effect of Cold Exposure on Determinants of Glucose Tolerance
To investigate mechanisms underlying the coordinate regulation of insulin sensitivity and insulin secretion, we sought to validate previous evidence that cold exposure inhibits insulin secretion in rats (16,19). As expected (16,26), fasting levels of blood glucose (112.6 ± 3.5 vs. 97.5 ± 2.3 mg/dL; P < 0.05) and plasma insulin (22.3 ± 2.9 vs. 10.0 ± 1.1 μU/mL; P < 0.05) were lower in fasted rats housed in a cold environment (5°C for 28 h) than at room temperature (22°C) (Fig. 2A and B). Moreover, in rats that underwent a FSIGT, followed by minimal model analysis (20,23,24,31), as expected, AIRG was potently inhibited by cold exposure by ~50% (AIRG: 667 ± 41 for room temperature vs. 372 ± 33 μU/mL × min for cold exposure; P < 0.001), despite glucose tolerance remaining virtually unchanged (AUCglucose: 5,869 ± 233 for room temperature vs. 5,848 ± 442 for cold exposure; P = NS) (Fig. 2A and B) (16). To explain this outcome, we hypothesized that a compensatory increase of insulin sensitivity must have occurred in the cold-exposed group. As predicted, we found that the potent effect of cold exposure to inhibit insulin secretion was offset by a near doubling of insulin sensitivity (SI) (Fig. 2C and D). Consequently, neither the DI nor glucose tolerance was substantially altered (Fig. 2E). In addition, cold exposure had no significant effect on SC, the BIE, or GEZI (Fig. 2F–H).

Reversibility of the Effect of Cold Exposure on Determinants of Glucose Tolerance
To assess the durability of the potent and highly coordinated (and offsetting) changes of insulin secretion and insulin sensitivity elicited by cold exposure, we performed an FSIGT on separate cohorts of fasted rats that were maintained at room temperature or were housed in the same cold environment (5°C) for 24 h and then returned to room temperature (22°C) for 4 h before study. Although fasting blood glucose and plasma insulin levels remained slightly lower in cold-exposed animals that were then returned to room temperature (Fig. 3A and B), the effect of cold exposure to inhibit glucose-induced insulin secretion (AIRG) (Fig. 3B) and/or increase SI (Fig. 2D) was no longer evident 4 h after returning to room temperature (Fig. 3D). As before, neither DI nor SC (Fig. 3E and F), GEZI, or BIE (data not shown) differed between groups (Fig. 3E). Thus, the potent and highly coordinated effects of cold exposure on insulin secretion and insulin sensitivity are reversed within 4 h after the return to room temperature.
Effect of Phentolamine on Determinants of Glucose Tolerance in Cold-Exposed Rats

To determine whether a change in SNS output contributes to the rapidly reversible effect of cold exposure on insulin secretion and insulin sensitivity, fasted rats housed at 5°C for 28 h received a continuous i.v. infusion of the α-AR blocker phentolamine or its vehicle for 30 min before and then during an FSIGT. As predicted, the effect of cold exposure to suppress basal- and glucose-induced insulin secretion was reversed by i.v. phentolamine, such that the low plasma insulin levels typical of animals housed in the cold were restored to values observed in animals housed at room temperature (13.9 ± 2.4 vs. 28.4 ± 1.9 μU/mL; P < 0.05).

Effect of Phentolamine on Determinants of Glucose Tolerance in Cold-Exposed Rats

To determine whether a change in SNS output contributes to the rapidly reversible effect of cold exposure on insulin secretion and insulin sensitivity, fasted rats housed at 5°C for 28 h received a continuous i.v. infusion of the α-AR blocker phentolamine or its vehicle for 30 min before and then during an FSIGT. As predicted, the effect of cold exposure to suppress basal- and glucose-induced insulin secretion was reversed by i.v. phentolamine, such that the low plasma insulin levels typical of animals housed in the cold were restored to values observed in animals housed at room temperature (13.9 ± 2.4 vs. 28.4 ± 1.9 μU/mL; P < 0.05). Despite this marked increase of insulin secretion, glucose tolerance was once again largely unaffected (AUCglucose: 5,151 ± 351 for vehicle vs. 5,078 ± 378 for phentolamine; P = NS) (Fig. 4B and C). In addition, the potent effect of cold exposure to increase Sg was also rapidly reversed by phentolamine-induced α-AR blockade, such that the DI did not change (Fig. 4D and E). Interestingly, Sg was also increased in phentolamine-treated animals (Fig. 4F), an effect due largely to an increase of GEZI with no change in BIE (data not shown). Collectively, these data show that the potent, highly coordinated, and reciprocal adjustments of insulin secretion and insulin sensitivity induced by cold exposure are both reversible within 30 min and dependent on intact α-AR signaling.

Effect of Phentolamine on Insulin Sensitivity Measured by Euglycemic-Hyperinsulinemic Clamp in Cold-Exposed Rats

To confirm our finding that α-AR blockade reverses the effect of cold exposure to increase insulin sensitivity, we measured the effect of i.v. phentolamine infusion on the glucose infusion rate (GIR), a direct measure of whole-body insulin sensitivity, in cold-exposed rats during a euglycemic-hyperinsulinemic clamp. By design, arterial blood glucose
levels were similar between groups during the clamp period (Fig. 5A), although plasma insulin levels during the clamp were significantly increased in cold-exposed animals receiving phentolamine (Fig. 5B), raising the possibility that phentolamine infusion affected insulin clearance as well as insulin secretion. Yet despite higher plasma insulin levels, the GIR required to maintain euglycemia was decreased by \( \frac{1}{4} \) \( (P < 0.05) \) in cold-exposed rats that received i.v. phentolamine (Fig. 5C and D). This finding offers direct, independent confirmation of the effect of systemic α-AR blockade to reduce insulin sensitivity in cold-exposed rats.

**Effects of Cold Exposure With or Without Phentolamine Infusion on Insulin Clearance From Plasma and on Other Humoral Determinants of Glucose Homeostasis**

To determine whether either cold exposure or phentolamine administration affected insulin clearance from plasma as well as insulin secretion, we measured C-peptide levels in plasma samples obtained during the FSIGT studies reported above and subjected these data to a model-based deconvolution analysis (25). We found that changes in the plasma level of C-peptide closely paralleled those of insulin from both studies (Fig. 6A and B). Further analysis also revealed no difference in rates of insulin clearance between animals housed at room temperature versus the cold (0.261 vs. 0.279; \( P = 0.59 \)) or between cold exposed animals treated with i.v. vehicle versus phentolamine (0.184 vs. 0.199; \( P = 0.31 \)). Both the effect of cold exposure to reduce the plasma insulin response to glucose and the effect of phentolamine to reverse this effect, therefore, resulted from changes of insulin secretion rather than insulin clearance.

An increase in nocturnal NEFA levels has been reported to contribute to the compensatory increase of insulin secretion in a dog model of insulin resistance induced by HFD feeding (8,9). However, we found no difference of fasting NEFA levels between rats housed at room temperature and cold-exposed animals. Further, NEFA levels tended to increase in cold-exposed animals during the FSIGT, whereas insulin secretion was reduced (Fig. 6C). Combined with the observation that plasma NEFA levels dropped comparably in the first 10 min after administration of glucose in cold-exposed animals, irrespective of whether they received i.v. saline or phentolamine (\( P = \text{NS} \)) (Fig. 6D), our data suggest that the effect of phentolamine to acutely
increase insulin secretion in cold-exposed rats was not a consequence of increased plasma NEFA levels.

In addition, we found that plasma glucagon levels were elevated in cold-exposed animals relative to those housed at room temperature (Fig. 6E) and were similarly suppressed after an i.v. glucose bolus during the FSIGT, irrespective of the temperature at which they were housed (Fig. 6F). Similarly, plasma corticosterone levels were elevated in cold-exposed rats, and phentolamine treatment, if anything, lowered plasma corticosterone levels (Fig. 6G and H), despite reversing the effect of cold to improve insulin sensitivity (Figs. 4 and 5). Taken together, these data suggest that neither changes in circulating NEFA levels nor the neuroendocrine response to cold is likely to explain the robust and highly coordinated changes of insulin secretion and insulin sensitivity.

**DISCUSSION**

That cold exposure elicits increased whole-body glucose utilization (to support the increased demand for heat production) (19) and decreased insulin secretion (16,32) with little or no change of glucose tolerance points to a highly coordinated metabolic response that enables thermogenic needs to be met while preserving glucose homeostasis. Our data suggest that these adaptive changes of insulin secretion and sensitivity to cold exposure occur quickly via a mechanism involving the SNS. Specifically, we found that the effect of cold exposure to inhibit the insulin response to glucose (whether measured as AIRG or AUCinsulin) in rats was precisely offset by a proportionate increase of insulin sensitivity (measured as $S_I$), such that glucose tolerance was unchanged. Moreover, this reduced insulin response to glucose was due entirely to reduced insulin secretion with no change of plasma insulin clearance, was fully established within 28 h of exposure to a cold environment, and returned to baseline within just 4 h after the return to room temperature. Our finding that the effect of cold exposure on these parameters was fully reversed by $\alpha$-AR blockade implicates the SNS in the coordinate regulation of insulin secretion and sensitivity in this setting. That this reversal was achieved within just 30 min of the onset of phentolamine infusion attests to the remarkable rapidity with which this coordinate regulation takes place.

The proper interpretation of these findings partly hinges on the extent to which the well-documented hyperbolic relationship between insulin secretion and insulin sensitivity

**Figure 4**—Role of the SNS on determinants of glucose tolerance in cold-exposed rats. Blood glucose levels (A), plasma insulin levels (B), AIRG (C), insulin sensitivity ($S_I$) (D), DI (E), and $S_G$ (F) in fasted adult male Wistar rats that were exposed to the cold at 4°C for 28 h and given a continuous i.v. infusion of the $\alpha$-AR blocker phentolamine (8 μg/kg/min) or vehicle, beginning 30 min before and continuing throughout a FSIGT. Mean ± SEM. *P < 0.05 vs. vehicle.
in humans (3) applies to rats as well. To address this question, we analyzed the relationship between $S_I$ and two different measures of the insulin response to a glucose challenge (AIRG and AUCinsulin) across a large cohort of normal rats housed at room temperature (22°C). To ensure a sufficiently broad distribution of these parameters, we included data from two different rat strains (Wistar and Long-Evans) fed either of two diets (standard chow or a 60% kcal HFD). As predicted, the relationship between $S_I$ and each of the two measures of insulin response was largely hyperbolic. Although we also found that SG makes a major contribution to glucose tolerance, which is consistent with prior evidence (33) and may help explain why the RMA regression hyperbolic fits included significant nonzero asymptotic values (intercept terms), our data establish that the relationship between insulin sensitivity and insulin secretion in rats is quite similar to that in humans. By recapitulating the relationship observed in humans, our data also support the validity of the FSIGT/minimal model method for metabolic studies in rats. Extending this validation is our findings that reliable exposure to cold reduces insulin sensitivity in otherwise normal rats and that this effect is offset by a proportionate increase of insulin secretion (Fig. 1C), and 2) in cold-exposed rats, the effect of i.v. phentolamine to rapidly reduce insulin sensitivity when measured as $S_I$ was replicated using the euglycemic-hyperinsulinemic clamp technique.

By investigating the role played by the brain (acting via the SNS) in the adaptive response to a physiological challenge, our approach is a departure from more conventional strategies that rely on administration of peptides or drugs into the brain to investigate its role in glucose homeostasis. Indeed, metabolic studies are usually conducted under constant environmental conditions to avoid the potentially confounding effect of commonly encountered environmental challenges. This strategy reduces variability in measured end points but also precludes investigation into how animals adapt to such challenges. Should the brain's role in glucose homeostasis take on greater importance when animals are confronted with a real-world environmental challenge, therefore, housing them in an unchanging (and unchallenging) environment by definition places major limits on the insights that can be gained. In this context, we note that the current study paradigm involved exposure to temperatures comparable to what rodents throughout much of the world experience daily.

To understand how cold exposure effects these highly coordinated changes of insulin secretion and insulin sensitivity, we consider three possibilities: 1) adaptive changes of insulin sensitivity are secondary to the change of insulin secretion, 2) adaptive changes of insulin secretion are secondary to the change of insulin sensitivity, or 3) adaptive changes of both insulin sensitivity and insulin secretion are part of an integrative regulatory process involving the SNS. With respect to the first possibility, we note previous evidence demonstrating that insulin sensitivity does not increase when insulin secretion is reduced, whether by a pancreatic β-cell toxin (e.g., streptozotocin) (34), inhibitory hormones (e.g., somatostatin) (35), partial
The alternative possibility that reduced insulin secretion is a consequence of a cold-induced increase of insulin sensitivity is consistent with the known effect of cold exposure to increase insulin-mediated glucose uptake into thermogenic tissues, including skeletal muscle, heart, and white and brown adipose tissue (17–19). One potential mechanism to explain how this effect is coupled to a proportionate decrease of insulin secretion is through reduced nutrient stimulation of β-cells via a mechanism involving, for example, reduced circulating glucose or NEFA levels (8–10). Such a mechanism is inconsistent with both the modest effect of cold exposure on plasma levels of these nutrients and the lack of any effect of phentolamine on these plasma values (despite its ability to fully reverse cold-induced inhibition of insulin secretion). Similarly, although cold exposure increased circulating levels of glucagon and corticosterone via a mechanism that was partially reversed by phentolamine, such changes do not offer a viable explanation for the associated changes of insulin secretion and sensitivity.
Could cold-induced changes of insulin secretion and insulin sensitivity be mediated via the SNS? That the effect of cold exposure to increase insulin sensitivity was reversed by $\alpha$-AR blockade offers direct evidence that increased SNS outflow is required for this effect, consistent with an extensive literature implicating the brain in the control of insulin sensitivity (38) via mechanisms involving the SNS (39). The hypothesis that the cold-induced decline of insulin secretion was also mediated via the SNS is supported by pioneering work documenting autonomic control of pancreatic islet function some 40 years ago (15) and by evidence that cold exposure increases sympathetic tone to the pancreas (12). The mechanism underlying this effect was subsequently shown to involve $\alpha$-AR signaling in pancreatic islets (13–15), a possibility reinforced by our finding that the effect of cold exposure to reduce insulin secretion was rapidly reversed (within 30 min) by phentolamine administration.

These observations collectively support a model in which SNS outflow to both pancreatic islets and thermogenic tissues underlies the highly coordinated and proportionate changes of insulin sensitivity and insulin secretion that effectively redirect substrate to thermogenic tissues without altering glucose tolerance (Fig. 7). Future studies are warranted to clarify the cellular basis for these effects and to further delineate the brain’s role in these metabolic responses. For example, the specific role played by islet sympathetic nerves could be investigated by sympathetic denervation of the pancreas or by targeted deletion of $\alpha$-AR in pancreatic $\beta$-cells. An additional priority is to determine whether reciprocal adjustment of insulin secretion and insulin sensitivity is triggered by activation of hypothalamic cold-sensitive neurons involved in thermoregulation.

In conclusion, we report that the hyperbolic relationship between insulin secretion and insulin sensitivity characteristic of humans is operational in rats as well. We demonstrate further that in rats, the rapid, potent, and highly coordinated changes of insulin sensitivity and insulin secretion elicited by cold exposure are critically dependent on the SNS. We speculate that these metabolic adjustments are components of a larger and highly integrated set of responses elicited by the brain to enable effective thermogenesis while preserving metabolic homeostasis and thereby promote survival during cold exposure.

Acknowledgments. The authors greatly acknowledge valuable guidance provided by Dr. David Wasserman, Vanderbilt University, and technical assistance provided by Amy Martinson, from the laboratory of Charles E. Murry, University of Washington, for measures of heart rate and mean arterial pressure.

Funding. This work was supported by National Institutes of Health, National Institute of Diabetes and Digestive and Kidney Diseases grants DK-089056 (G.J.M.), DK-27619 and DK-29867 (R.N.B.), DK-50154 (G.J.T.), DK-083042, DK-090320, and DK-101997 (M.W.S.); the National Institute of Diabetes and Digestive Kidney Diseases–funded Nutrition Obesity Research Center (DK-035816) and Diabetes Research Center (DK-017047); the Nutrition, Obesity and Atherosclerosis Training Grant from the National Heart, Lung, and Blood Institute (T32-HL-007028) and the Diabetes and Metabolism Training Grant (T32-DK-0007247) at the University of Washington; and the Department of Veterans Affairs grant BX001060 (S.E.K.).

Duality of Interest. No potential conflicts of interest relevant to this article were reported.

Author Contributions. G.J.M., K.M., K.J.K., F.P., D.S., R.N.B., G.J.T., S.E.K., and M.W.S. analyzed data and edited the manuscript. G.J.M., K.M., J.M.R., J.M.S., M.E.M., J.T.N., and N.K.A. conducted experiments and acquired data. G.J.M., K.M., and M.W.S. designed research studies and wrote the manuscript. M.W.S. is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.
References

1. Bergman RN, Ider YZ, Bowden CR, Cobelli C. Quantitative estimation of insulin sensitivity. Am J Physiol 1979;236:E667–E677
2. Bergman RN, Phillips LS, Cobelli C. Physiologic evaluation of factors controlling glucose tolerance in man: measurement of insulin sensitivity and beta-cell glucose sensitivity from the response to intravenous glucose. J Clin Invest 1981;68:1456–1467
3. Kahn SE, Prigeon RL, McCulloch DK, et al. Quantification of the relationship between insulin sensitivity and beta-cell function in human subjects. Evidence for a hyperbolic function. Diabetes 1993;42:1663–1672
4. Buchanan TA, Metzger BE, Freinkel N, Bergman RN. Insulin sensitivity and B-cell responsiveness to glucose during late pregnancy in lean and moderately obese women with normal glucose tolerance or mild gestational diabetes. Am J Obstet Gynecol 1990;162:1008–1014
5. Moran A, Jacobs DR Jr., Steinberger J, et al. Insulin resistance during puberty: results from clamp studies in 357 children. Diabetes 1999;48:2039–2044
6. Kahn SE, Hull RL, Utzschneider KM. Mechanisms linking obesity to insulin resistance and type 2 diabetes. Nature 2006;444:840–846
7. Lorenzo C, Wagenknecht LE, Rewers MJ, et al. Disposition index, glucose effectiveness, and conversion to type 2 diabetes: the Insulin Resistance Atherosclerosis Study (IRAS). Diabetes Care 2010;33:2098–2103
8. Kim SP, Catalano KJ, Hsu IR, Chiu JD, Richy J, Bergman RN. Nocturnal free fatty acids are uniquely elevated in the longitudinal development of diet-induced insulin resistance and hyperinsulinemia. Am J Physiol Endocrinol Metab 2007;292:E1590–E1598
9. Broussard JL, Kolka CM, Castro AV, et al. Elevated nocturnal NEFA are an early signal for hyperinsulinemic compensation during diet-induced insulin resistance in dogs. Diabetologia 2015;58:2663–2670
10. Maickel RP, Matussek N, Stern DN, Brodie BB. The sympathetic nervous system as a homeostatic mechanism. I. Absolute need for sympathetic nervous function in body temperature maintenance of cold-exposed rats. J Pharmacol Exp Ther 1967;157:103–110
11. Morrison SF, Madden CJ, Tupone D. Central neural regulation of brown adipose tissue thermogenesis and energy expenditure. Cell Metab 2014;19:741–756
12. Young JB, Landsberg L. Effect of diet and cold exposure on norepinephrine turnover in pancreas and liver. Am J Physiol 1979;236:E524–E533
13. Ahrén B, Taborsky GJ Jr., Porter D Jr. Neuropeptidergic versus cholinergic adrenergic regulation of islet hormone secretion. Diabetes Metab Rev 1986;29:827–836
14. Fagerholm V, Haaparanta M, Scheinin M. α2-adrenoceptor regulation of blood glucose homeostasis. Basic Clin Pharmacol Toxicol 2011;108:365–370
15. Porte D Jr., Robertson RP. Control of insulin secretion by catecholamines, stress, and the sympathetic nervous system. Fed Proc 1973;32:1792–1796
16. Vallerand AL, Lupien J, Bukowiecki LJ. Interactions of cold exposure and starvation on glucose tolerance and insulin response. Am J Physiol 1983;245: E575–E581
17. Vallerand AL, Pérusse F, Bukowiecki LJ. Stimulatory effects of cold exposure and cold acclimation on glucose uptake in rat peripheral tissues. Am J Physiol 1990;259:R1043–R1049
18. Smith SA, Young P, Cawthorne MA. Quantification in vivo of the effects of insulin on glucose utilization in individual tissues of warm- and cold-acclimated rats. Biochem J 1986;237:789–795
19. Vallerand AL, Pérusse F, Bukowiecki LJ. Cold exposure potentiates the effect of insulin on in vivo glucose uptake. Am J Physiol 1987;253:E179–E186
20. Rojas JM, Matsen ME, Mundinger TO, et al. Glucose intolerance induced by blockade of central FGF receptors is linked to an acute stress response. Mol Metab 2015;4:561–568
21. German J, Kim F, Schwartz GJ, et al. Hypothalamic leptin signaling regulates hepatic insulin sensitivity via a neurocircuit involving the vagus nerve. Endocrinology 2009;150:4502–4511
22. German JP, Thaler JP, Wisse BE, et al. Leptin activates a novel CNS mechanism for insulin-independent normalization of severe diabetic hyperglycemia. Endocrinology 2011;152:394–404
23. Morton GJ, Matsen ME, Brady DP, et al. FGF19 action in the brain induces insulin-independent glucose lowering. J Clin Invest 2013;123:4799–4808
24. Alonso LC, Watanabe Y, Stefanovski D, et al. Simultaneous measurement of insulin sensitivity, insulin secretion, and the disposition index in conscious unhandled mice. Obesity (Silver Spring) 2012;20:1403–1412
25. Ahrén B, Thomasset K, Pacini G. Reduced insulin clearance contributes to the increased insulin levels after administration of glucagon-like peptide 1 in mice. Diabetologia 2005;48:2140–2146
26. Beck LV, Zaharko DS, Kalsec SR. Variation in serum insulin and glucose of rats with chronic cold exposure. Life Sci 1967;6:1501–1506
27. Bergman RN. Minimal model: perspective from 2005. Horm Res 2005;64 (Suppl. 3):8–15
28. Sokal RR, Rohlf FJ. Biometry: The Principles and Practice of Statistics in Biological Research. 4th ed. New York, W.H. Freeman, 2012
29. Boston RC, Moate PJ, Stefanovski D, Sumner AE, Bergman RN. AKA-glucose: a program for kinetic and epidemiological analysis of frequently sampled intravenous glucose tolerance test data using database technology. Diabetes Technol Ther 2005;7:298–307
30. Kahn SE, Prigeon RL, McCulloch DK, et al. The contribution of insulin-dependent and insulin-independent glucose uptake to intravenous glucose tolerance in healthy human subjects. Diabetes 1994;43:587–592
31. Best JD, Kahn SE, Ader M, Watanabe RM, Ni TC, Bergman RN. Role of glucose effectiveness in the determination of glucose tolerance. Diabetes Care 1996;19:1018–1030
32. Vallerand AL, Frim J, Kavanagh MF. Plasma glucose and insulin responses to oral and intravenous glucose in cold-exposed humans. J Appl Physiol (1985) 1988;65:2395–2399
33. Bunner AE, Chandrasekera PC, Barnard ND. Knockout mouse models of insulin signaling: relevance past and future. World J Diabetes 2014;5:146–159
34. Tobin BL, Finegood DT. Reduced insulin secretion by repeated low doses of STZ impairs glucose effectiveness but does not induce insulin resistance in dogs. Diabetes 1993;42:474–483
35. Kahn SE, Klaff LJ, Schwartz MW, et al. Treatment with a somatostatin analog decreases pancreatic B-cell and whole body sensitivity to glucose. J Clin Endocrinol Metab 1990;71:994–1002
36. Ward WK, Wallum BJ, Beard JC, Taborsky GJ Jr., Porte D Jr. Reduction of glycemic potentiation. Sensitive indicator of beta-cell loss in partially pancreatectomized dogs. Diabetes 1988;37:723–729
37. Utzschneider KM, Prigeon RL, Carr DB, et al. Impact of differences in fasting glucose and glucose tolerance on the hyperbolic relationship between insulin sensitivity and insulin responses. Diabetes Care 2006;29:356–362
38. Schwartz MW, Seeley RJ, Tschop MH, et al. Cooperation between brain and islet in glucose homeostasis and diabetes. Nature 2013;503:59–66
39. Haque MS, Minokoshi Y, Hamai M, Iwai M, Horiiuchi M, Shimizu T. Role of the sympathetic nervous system and insulin in enhancing glucose uptake in peripheral tissues after intrahypothalamic injection of leptin in rats. Diabetes 1999;48:1706–1712