INTRODUCTION

Glucose is the primary energy source for cellular metabolism. Therefore, the maintenance of appropriate serum glucose levels is critical for organismal survival. Glucose homeostasis is maintained through interplay between central and peripheral control mechanisms which are aimed at storing excess glucose following meals and mobilizing these same stores during periods of fasting. Central control of glucose homeostasis is accomplished by a distributed system of glucosensing elements, integrative neural networks which process signals from these elements, and pre-autonomic nuclei which control motor outflow and are modulated through inputs from these neural integrators (Watts and Donovan, 2010).

A principal site for neural integration of glucose availability is the nucleus of the solitary tract (NST), located in the dorsal medulla. Cells within NST have unique access to serum glucose levels due to the large numbers of fenestrated capillaries present in the nucleus (Gross et al., 1990) as well as its close apposition to the area postrema, a circumventricular organ (Cottrell and Ferguson, 2004). A subset of NST neurons have been shown to respond to physiologically relevant changes in extracellular glucose availability both in vivo (Yettefti et al., 1995, 1997) and in vitro (Mizuno and Oomura, 1984; Himmi et al., 1996; Dallaporta et al., 1999). In addition to containing these glucosensitive cells, NST also receives input regarding peripheral glucose levels by way of glucosensors located in hepatic portal/mesenteric vein, which innervate NST via the hepatic branch of the vagus nerve (Niijima, 1984; Adachi et al., 1995; Grabauskas et al., 2010). Furthermore, NST also receives descending inputs from hypothalamic nuclei such as the lateral and paraventricular nuclei that are associated with glucose homeostasis (Marty et al., 2007; Geerling et al., 2010; Biag et al., 2012). Due to this confluence of signals regarding glucose availability as well as its inputs to pre-autonomic nuclei, the NST plays a predominant role in central control of glucose homeostasis (Adachi et al., 1995; Grill and Hayes, 2012).

The NST is also commonly associated with the physiological response to hypoglycemia. These autonomic counter-measures, often referred to collectively as the counter-regulatory response (CRR) include increases in serum levels of glucagon and stress hormones, increased food intake, and an overall increase in sympathetic tone. Systemic administration of the glucoprivic compound 2-deoxy-glucose (2-DG; a glucose analog commonly used to invoke the CRR) results in increased c-Fos expression in the NST (Ritter et al., 1998; Briski and Marshall, 2000; Sanders and Ritter, 2000; Dodd et al., 2010). Focal administration of a similar glucoprivic agent, 5-thio-D-glucose, directly into the NST of rats also drives CRRs such as increased food intake (Ritter et al., 2000) and increases in serum glucagon and stress hormone levels (Andrew et al., 2007).

Glucose homeostasis is maintained through interplay between central and peripheral control mechanisms which are aimed at storing excess glucose following meals and mobilizing these same stores during periods of fasting. The nucleus of the solitary tract (NST) in the dorsal medulla has long been associated with the central detection of glucose availability and the control of glucose homeostasis. Recent evidence has emerged which supports the involvement of astrocytes in glucose homeostasis. The aim of the present study was to investigate whether NST-astrocytes respond to physiologically relevant decreases in glucose availability, in vitro, as well as to the presence of the glucoprivic compound 2-deoxy-D-Glucose. This report demonstrates that some NST-astrocytes are capable of responding to low glucose or glucoprivation by increasing cytoplasmic calcium; a change that reverses with restoration of normal glucose availability. While some NST-neurons also demonstrate an increase in calcium signaling during low glucose availability, this effect is smaller and somewhat delayed compared to those observed in adjacent astrocytes. TTX did not abolish these hypoglycemia mediated responses of astrocytes, suggesting that NST-astrocytes may be directly sensing low glucose levels as opposed to responding to neuronal detection of hypoglycemia. Thus, chemodetection of low glucose by NST-astrocytes may play an important role in the autonomic regulation of glucose homeostasis.

Keywords: astrocytes, calcium imaging, 2-deoxy-D-glucose, solitary nucleus, vago-vagal reflex, glucose homeostasis, hypoglycemia
In addition to its role in glucose homeostasis and hypoglycemic counter-regulation, the NST plays a critical role in a number of other autonomic functions such as regulation of cardiovascular, respiratory, and gastrointestinal reflexes (Blessing, 1997). There is increasing evidence that chemodetection by brainstem astrocytes can drive changes in these autonomic reflexes (Hermann and Rogers, 2009; Hermann et al., 2009; Gourine et al., 2010; Kasymov et al., 2013). Interestingly there is also strong evidence for astrocytic involvement in the autonomic response to hypoglycemia. Systemic administration of the selective glial toxin, methionine sulfoximine, blocks 2-DG induced c-Fos expression in NST (Young et al., 2000). In addition, transgenic mice which only express the type II glucose transporter (GLUT2) in pancreatic beta cells (i.e., expression of GLUT2 was knocked out centrally), demonstrate defects in the CRR (Burcelin and Thorens, 2001). However, these defects are rescued by the centrally expressed beta cells (i.e., expression of GLUT2 was knocked out in vivo), demonstrating defects in the CRR (Burcelin and Thorens, 2001). Therefore, the aim of the present study was to investigate whether NST-astrocytes respond to physiologically relevant decreases in glucose availability, in vitro, as well as to the presence of the glucoprivic compound 2-DG.

**METHODS**

**ANIMALS**

All experimental procedures were conducted under the approval of the Pennington Biomedical Research Center’s Institutional Animal Care and Use Committee. Long Evans rats of either sex (body weight between 150 and 250 g) were used in these studies. All animals were housed in a temperature controlled room under 12 h light/dark cycle and provided water and food ad libitum. All experimental protocols were performed according to the guidelines set forth by the National Institutes of Health.

**In vivo PRE-LABELING OF NST AND HARVEST OF HINDBRAIN SLICES**

Animals (N = 22) were deeply anesthetized with urethane (1.5 g/kg, ip; ethyl carbamate, Sigma) and placed in a stereotaxic frame. Using aseptic technique, the occipital plate of the skull was removed to expose the medullary brainstem. A micropipette 30 micron tip diameter; filled with 0.2% Calcium Green 1 AM (CAG; Life Technologies), 0.3% sulforhodamine 101 (SR101; Sigma Chemical) and 20% pluronic-DMSO (F-127, in pH 7.2 tris-PBS buffer) was directed toward the medial solitary nucleus using a stereotaxic carrier. Four injections (40 nL each) of the CAG-SR101 solution were made unilaterally into the NST at the level of calamus and 0.2, 0.4, and 0.6 mm anterior to calamus; all at a depth of 300 microns below the surface. This injection pattern labeled the entire ipsilateral medial NST (Hermann et al., 2009). CAG, a calcium reporter dye, is taken up by both neurons and glia while SR101 labels only astrocytes (Nimmerjahn et al., 2004; McDougal et al., 2011). SR101 does not interfere with CAG fluorescence (Hermann and Rogers, 2009). Thus, despite the similarity in size of NST neuronal and astrocytic cell bodies, this method made it possible to easily discriminate between these two cell types in the slice preparation (Figure 1). After a 30 min interval to allow for dye uptake, the anesthetized rat was decapitated and the brainstem was quickly harvested. The caudal brainstem was glued to an aluminum block and placed in cold (~4°C) carboxygenated (95% O2; 5% CO2) cutting solution (recipe below). Coronal sections (300 micron thick) were cut through the medulla using a Leica VT1200 tissue slicer equipped with a sapphire knife (Delaware Diamond Knives). The NST is easily identified in these slices due to its lack of myelin; 4–5 slices containing NST were collected from each animal and placed in normal Krebs’ solution (recipe below) which was bubbled with 95% O2/5% CO2 and maintained at a constant temperature of 29°C. Slices were allowed to equilibrate to these conditions for an hour prior to imaging.

**In vitro DRUGS AND SOLUTIONS**

All solutions were freshly prepared on the day of the experiment. The cutting solution contained 110 mM choline chloride, 25 mM NaHCO3, 2.5 mM KCl, 7 mM MgSO4–7H2O, 1.25 mM NaH2PO4, 10 mM glucose, and 0.5 mM CaCl2–2H2O. The normal Krebs’ solution contained 124 mM NaCl, 25 mM NaHCO3, 3.0 mM KCl, 1 mM MgSO4–7H2O, 1.5 mM NaH2PO4, 5 mM glucose, and 1.5 mM CaCl2–2H2O. Composition of the “low glucose challenge” solutions was identical to the normal Krebs’ solution except that the concentration of glucose was decreased from 5 to 2.5 mM, 1 mM, or 0.5 mM. Given that extracellular brain glucose levels are ~30% that of plasma glucose (Silver and Erecinska, 1994), the range of normal extracellular brain glucose levels runs between 0.5 and 2.5 mM (Song et al., 2001). However, brain areas such as the arcuate nucleus of the hypothalamus or the NST, which border circumventricular organs, are predicted to be exposed to glucose levels normally seen in the CSF (3.4 mM) or serum (5.5 mM) (Levin et al., 2004). Therefore, we include 5 mM glucose as our “normal” glucose concentration for comparisons. The “glucoprivic challenge” solution consisted of 5 mM 2-deoxy-D-glucose (2-DG; Sigma-Aldrich, St. Louis, MO) dissolved in normal 5 mM glucose Krebs’ solution. The osmolality of
the normal Krebs', low glucose challenge, and glucoprivic chal-
lenge solutions were adjusted to 300 ± 3 mOsm with sucrose.
Tetrodotoxin (1 µM TTX; Sigma-Aldrich, St Louis, MO) was
included in some low glucose challenge solutions. Bath applica-
tion of TTX has been shown to block approximately 95% of the
direct neuronal activation of NST astrocytes (McDougal et al.,
2011).

**LIVE CELL CALCIUM IMAGING**
Slices were transferred to a custom imaging chamber (Rogers and
Hermann, 2012b); perfused at a rate of 2.5 mL/min with normal
Krebs’ maintained at a temperature of 32°C. Slices were imaged
with a Nikon F1 fixed stage upright microscope equipped with a
Nikon Fast Scan laser confocal head and a Luca EMCCD camera
(Andor Technology, South Windsor, CT). Co-injection of CAG
and SR101 allows for discrimination of astrocytes and neurons at
the cellular level by comparing images captured using the 488 and
561 nm laser lines (McDougal et al., 2011). That is, at 488 nm,
both astrocytes and neurons pre-labeled with CAG will appear
green; at 561 nm, all astrocytes will also appear red. A single,
dual exposure image was collected just prior to and following
each experimental trial in order to confirm the cell types being
recorded (Figure 1).

Changes in intracellular calcium concentrations within CAG
pre-labeled NST-astrocytes and NST-neurons in response to dif-
ferent stimuli were recorded using the 488 nm laser line to
excite the CAG. Increases in intracellular calcium concentra-
tions are reflected as an increase in fluorescence and is indicative
of increased cellular activity. During experimental trials, time-
lapsed images of mixed fields of NST-astrocytes and neurons
were monitored for their responses to low glucose or glucoprivic
conditions.

**EXPERIMENTAL DESIGN**
Each slice was exposed to a single “experimental trial” which con-
sisted of three perfusion conditions of equal duration (i.e., 10 min
for each condition). Slices were perfused with normal Krebs’ con-
taining 5 mM glucose during the initial and final blocks of time.
The intervening block was the experimental condition [e.g., “time
control” consisting of continued perfusion with 5 mM glucose
Krebs’, low glucose (i.e., 0.5, 1.0, or 2.5 mM) or glucoprivic chal-
genue (i.e., 5 mM 2-DG with 5 mM glucose Krebs’)]. Thus, each
cell functioned as its own control and transitional differences in
activity due to changes in available glucose (i.e., decreasing or
increasing) could also be monitored.

In order to rule out the possibility that neuronal synaptic
input to glia was responsible for the astrocytic responses to low-
glucose conditions, tetrodotoxin (TTX) was used to block action
potential-mediated communication between neurons and astro-
cytes (McDougal et al., 2011). For this study, we specifically chose
the 1 mM glucose challenge, to ensure that the neurons would not
be compromised by very low glucose. In these experiments, TTX
(1 µM) was present during all three perfusion periods [i.e., nor-
mal (5 mM), low glucose challenge (1 mM), and normal (5 mM)
glucose]. The responses of cells during the low glucose challenge
plus TTX were then compared to responses of cells exposed to an
identical experimental trial that did not use TTX, i.e., identical
blocks of 0.5, 1 and 5 mM perfusion conditions without TTX.

Time-lapse images were analyzed using Nikon NIS-Elements
software. Each cell in a field was delineated as a region of inter-
est (ROI) and the average intensity of all pixels within each ROI
was calculated for each time point (Figure 1). These data were
exported to Microsoft Excel for analysis offline.

Relative changes in intracellular calcium in response to all
stimulation paradigms were quantified as percent changes in
fluorescence: \(\frac{\Delta F}{F}\)%, where \(F\) is the baseline fluorescence
intensity within an area of interest (e.g., the outline of an SR101-
labeled astrocyte) before stimulation, and \(\Delta F\) is the change
from this value resulting from stimulus-induced cellular activity
(Helmchen, 2000). Background fluorescence (i.e., non-responsive
areas in same field) was subtracted from both \(F\) and \(\Delta F\). An
individual cell was considered to have responded to a given stim-
ulus if the stimulus induced a peak change in \(\frac{\Delta F}{F}\) > 5%
(O’Malley et al., 2006; Rogers et al., 2006a,b; Hermann et al., 2009;
McDougal et al., 2011).

Parameters including average baseline fluorescence intensity,
peak intensity, as well as the onset of the response to the low
glucose or glucoprivic challenge were determined for each ROI.
These values were used to calculate: (a) latency to onset of
response, (b) peak response magnitude (percent increase in inten-
sity from baseline), (c) response offset (time at which the response
magnitude decayed to 1/e of the peak value: \(\sim 37\%\)), and (d) (response
magnitude (area under the curve from response onset to response
offset) for each ROI. In order to verify the macro-
defined values, each was displayed graphically on a line plot of
intensity vs. time and visually inspected for accuracy.

The criteria of low-glucose (or 2DG) responder types:

- **Positive**: Peak response > 5% and AUC > 175 (\(\frac{\Delta F}{F}\))% minute
  and \(\frac{\Delta F}{F}\) values returned to 1/e of peak (\(\sim 37\%\)) before the end
  of the trial.
- **Negative**: Peak response < -5% and AUC < -175 (\(\frac{\Delta F}{F}\))% minute
- **Sustained activation**: Peak response > 5% and \(\frac{\Delta F}{F}\) values
  failed to return to 1/e of peak (\(\sim 37\%\)) before the end of
  the trial. AUC is undefined in these cells because there is no
  “stop time” which is defined as the time at which \(\Delta F\) values
  returned to 1/e of peak (\(\sim 37\%\)).
- **Non-responders**: Any cell which did not meet the above criteria.

**DATA ANALYSIS**
The cell type (i.e., astrocyte or neuron) represented by each
ROI was determined by the presence or absence of SR101 stain-
ing. The average values of three separate response parameters
peak, latency, and response magnitude) were determined for
each cell type in each experimental condition, and then com-
pared statistically. Astrocytic response parameters across the three
concentrations of glucose were compared to neuronal response
parameters using two-way analysis of variance (i.e., cell type vs.
concentration of glucose). Separate analyses were made of the
average response parameters of astrocytes vs. neurons to a glu-
coprivic condition (i.e., 5 mM glucose plus 5 mM 2-DG) using
unpaired t-tests.
Studies using TTX to determine if astrocytic responses were secondary to neuronal activation used unpaired t-tests to evaluate cellular response parameters of astrocytes or neurons during the 1 mM glucose challenge with vs. without TTX in the perfusion solution. All statistical analyses were performed using GraphPad Prism software (GraphPad Software, La Jolla, CA); p-values < 0.05 were considered statistically significant.

RESULTS
A SUBSET OF NST-ASTROCYTES AND NEURONS RESPOND TO LOW GLUCOSE CHALLENGE

Brainstem slices challenged by conditions of low glucose availability (i.e., bath application of Krebs’ solution containing 0.5, 1.0, or 2.5 mM glucose) showed fluxes in intracellular calcium levels in both NST-astrocytes and neurons. The cellular responses were measured by changes in the fluorescent intensity of CAG relative to individual baseline levels. The relative changes in cytoplasmic calcium were expressed as percent changes in fluorescence \[\text{[(}\Delta F/F\text{)]}\%\] of the CAG reporter dye, where \(F\) is the intensity of the baseline fluorescence signal before stimulation, and \(\Delta F\) is the difference between the peak fluorescence intensity and the baseline signal (Figures 2, 4, 5).

Both NST-astrocytes and neurons displayed one of four distinct response patterns as a result of exposure to each low glucose challenge (Table 1). The most prevalent response pattern was the “non-responder” which showed no response to conditions of low glucose availability (47% of astrocytes; 51% of neurons). The second largest category encountered was a “low glucose positive” response pattern. Table 1 illustrates the classification of cellular response patterns to low glucose (top) or glucoprivic (bottom) challenges. Two-Way ANOVA: (C) Peak response and (D) response magnitude demonstrated significant main effects attributed to the cell type \((p < 0.05)\) as well as significant main effects attributed to the concentration of glucose \((p < 0.05)\) (see also Figure 3). These main effects are qualified due to the significant interactions between cell types and glucose concentrations. Thus, NST-astrocytes were more responsive than NST-neurons to low glucose conditions particularly at the 0.5 mM concentration. (E) Latency to onset of response to low glucose also demonstrated a significant main effect attributed to cell type \((p < 0.05)\); astrocytes responded at \(1.7 \pm 0.2 \text{ min}\) (mean ± s.e.m.) compared to neurons at \(2.6 \pm 0.2 \text{ min}\). * \(p < 0.05\).

|                   | Low glucose positive (%) | Low glucose negative (%) | Low glucose sustained (%) | Low glucose NR (%) |
|-------------------|--------------------------|--------------------------|--------------------------|-------------------|
| **Astrocytes**    |                          |                          |                          |                   |
| 263 total         | 39                       | 13                       | 1                        | 47                |
| **Neurons**       |                          |                          |                          |                   |
| 160 total         | 37                       | 11                       | 1                        | 51                |

|                   | 2DG positive (%) | 2DG negative (%) | 2DG sustained (%) | 2DG NR (%) |
|-------------------|-----------------|-----------------|-----------------|----------|
| **Astrocytes**    |                 |                 |                 |          |
| 118 total         | 42              | 8               | 3               | 47       |
| **Neurons**       |                 |                 |                 |          |
| 81 total          | 26              | 16              | 6               | 52       |

FIGURE 2 | Example of response patterns of low glucose-positive astrocytes and neurons in the hindbrain slice preparation to low glucose challenge. As seen in Figure 1, mixed fields of NST-astrocytes and neurons were recorded using confocal live cell calcium imaging. Changes in intracellular calcium levels were reflected as changes in intensity of the fluorescent calcium indicator dye, Calcium Green 1-AM. Both low glucose-positive (A) astrocytes and (B) neurons were activated in response to a change in the extracellular glucose concentration from 5 to 0.5 mM; this activation returned toward baseline levels following return to 5 mM glucose. [Traces in (A,B) are the average response of five representative low glucose positive astrocytes and neurons, respectively]. Two-Way ANOVA: (C) Peak response and (D) response magnitude demonstrated significant main effects attributed to the cell type \((p < 0.05)\) as well as significant main effects attributed to the concentration of glucose \((p < 0.05)\) (see also Figure 3). These main effects are qualified due to the significant interactions between cell types and glucose concentrations. Thus, NST-astrocytes were more responsive than NST-neurons to low glucose conditions particularly at the 0.5 mM concentration. (E) Latency to onset of response to low glucose also demonstrated a significant main effect attributed to cell type \((p < 0.05)\); astrocytes responded at \(1.7 \pm 0.2 \text{ min}\) (mean ± s.e.m.) compared to neurons at \(2.6 \pm 0.2 \text{ min}\). * \(p < 0.05\).
response pattern where cells showed an increase in intracellular calcium concentrations when exposed to low glucose and baseline levels of intracellular calcium concentrations returned following the return to 5 mM glucose Krebs’ solution (39% of astrocytes; 37% of neurons) (Figures 2A,B; Table 1). The remaining cells responded either with a decrease in intracellular calcium levels and were characterized as “low glucose negative” (13% of astrocytes; 11% of neurons) or displayed an increase in intracellular calcium signaling which failed to return to baseline levels following the return to 5 mM glucose levels. This last group is referred to as “sustained activation” (1% of astrocytes; 1% of neurons; Table 1).

RESPONSE TO LOW GLUCOSE CHALLENGE DIFFERED BASED ON CELL TYPE AND GLUCOSE CONCENTRATION

Three response parameters were measured in “low glucose positive” astrocytes and neurons following exposure to each low glucose challenge condition (0.5, 1.0, or 2.5 mM); see Figure 3 and Table 2. These parameters included: (1) response latency (time between onset of challenge and response onset), (2) response magnitude (area under the curve from response onset to response offset), and (3) peak response (percent increase in CAG intensity from baseline). Two-Way ANOVA revealed that latency to onset of response demonstrated a significant main effect attributed to cell type [$F(1, 111) = 6.38, p < 0.05$]. Latency to onset of response to low glucose for astrocytes was 1.7 ± 0.2 min (mean ± SEM) compared to neurons at 2.6 ± 0.2 min. Peak response and response magnitude also demonstrated significant main effects attributed to the cell type [$F(1, 159) = 8.36$ and 12.57, respectively; $p < 0.05$] as well as significant main effects attributed to the concentration of glucose [$F(3, 159) = 8.40$ and 18.57; respectively; $p < 0.05$]. These main effects are qualified due to the significant interactions between cell types and glucose concentrations [$F(3, 159) = 5.24$ and 4.99, respectively; $p < 0.05$]. Thus, astrocytes were more responsive than neurons to low glucose conditions particularly at the 0.5 mM concentration (Figure 3).

Bonferroni post-hoc t-tests showed that astrocytes and neurons differ in these response parameters to 0.5 mM glucose in the perfusion bath (Figure 2). Additionally, astrocytes demonstrated significantly larger peak responses and response magnitudes to 0.5 mM glucose compared to either 1.0 or 2.5 mM glucose challenges (Figure 3).

A SUBSET OF NST-ASTROCYTES AND NEURONS RESPOND TO GLUCOPRIVIC CHALLENGE

NST-astrocytes and neurons showed similar response patterns during glucoprivic challenge where 5 mM 2-DG was included in the 5 mM glucose-Krebs’ solution (Figure 4). Most astrocytes (47%) and neurons (52%) were “non-responders” (Table 1). The

### Table 2 | Response parameters of NST astrocytes and neurons responsive to low-glucose or 2DG conditions.

| Condition     | Peak response(DF/F) % | Magnitude of response (Area under the curve = change over time) | N |
|---------------|-----------------------|---------------------------------------------------------------|---|
|               | Astrocytes            | Neurons            | Astrocytes | Neurons | Astrocytes | Neurons |
| 5 mM 2DG      | 15.5 ± 1.1            | 14.0 ± 1.0         | 2329 ± 228 | 2093 ± 229 | 56          | 21       |
| 0.5 mM glucose| 21.4 ± 1.8            | 10.4 ± 1.1         | 3005 ± 308 | 1224 ± 140 | 46          | 33       |
| 1.0 mM glucose| 10.1 ± 1.1            | 9.3 ± 1.8          | 678 ± 124  | 391 ± 60   | 35          | 17       |
| 2.5 mM glucose| 8.4 ± 0.7             | 7.8 ± 1.5          | 704 ± 139  | 391 ± 94   | 29          | 12       |

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FIGURE 4 | Activated response patterns of astrocytes and neurons in the hindbrain slice preparation to glucoprivic challenge (i.e., 2-DG-positive). Approximately 40% of NST-astrocytes and 25% of NST-neurons responded to a glucoprivic solution consisting of 5 mM 2-Deoxy-D-glucose (2-DG) dissolved in media containing normal (5 mM) glucose concentrations with an increase in activity (Tables 1, 2). Examples of 2-DG-positive astrocytes and neurons activated in response to the glucoprivic challenge are shown [Traces in (A,B) are the average response of five representative 2-DG-positive astrocytes and neurons, respectively]. The peak responses (C) and response magnitudes (D) were similar in both astrocytes and neurons. However, these two cell types differed in time to onset of response (E). NST-astrocytes averaged 1.4 ± 0.1 min compared to the onset of NST-neuronal response at 2.2 ± 0.2 min (*p < 0.05).

next largest class was “2-DG positive” which displayed an increase in intracellular calcium signaling during exposure to this glucoprivic condition that returned to baseline following washout (42% of astrocytes; 26% of neurons). The remainder of cells could be characterized as either “2-DG negative” (8% of astrocytes; 16% of neurons) or “sustained activation” (3% of astrocytes; 6% of neurons). Unpaired t-test comparisons revealed that astrocytes had significantly shorter latencies to response onset compared to neurons [1.4 ± 0.1 min vs. 2.2 ± 0.2 min; t(73) = 3.00; p = 0.0037]. There were no differences in peak response (15.5 ± 1.1% vs. 14.0 ± 1.0%; astrocytes vs. neurons) or response magnitude (2329 ± 228 vs. 2093 ± 229) between astrocytes and neurons during glucoprivic challenge (Table 2).

LOW GLUCOSE SIGNALING IN NST-ASTROCYTES IS INDEPENDENT OF NEURONAL SIGNALING
In order to rule out the possibility that neuronal synaptic input to glia was responsible for the astrocytic responses to low-glucose conditions, TTX was used to block action potential-mediated communication between neurons and astrocytes (Figure 5). Astrocytic peak responsiveness to a 1mM low glucose challenge was unaffected by TTX (15.3 ± 0.9% vs. 17.5 ± 1.7%), although response magnitude was somewhat lower than control levels (991 ± 102 vs. 763 ± 116; Figures 5A,C,D). The slight reduction in the size of the astrocytic responses in the presence of TTX likely represents a loss of tonic neuronal activation of NST-astrocytes independent of glucose availability, possibly mediated by glutamate release from vagal afferents (McDougal et al., 2011).

In contrast, neuronal peak response and response magnitude during the same low glucose challenge were significantly reduced by TTX [peak response: 9.0 ± 1.0% vs. 6.2 ± 0.6%; t(17) = 2.422, p = 0.013; magnitude: 652 ± 135 vs. 156 ± 37; t(17) = 3.043, p = 0.004 (Figures 5B,E,F)]. NST neuronal calcium signaling in response to low glucose challenge may be dependent on other neuronal inputs or on modulation of the voltage-dependent sodium conductance.

DISCUSSION
The dorsal medulla and NST have long been associated with the central detection of the glucose availability and the control of the glycemic state (Bernard, 1855; Marty et al., 2007; Grill and Hayes, 2012). Recent evidence is emerging in support of astrocytic involvement in glucose homeostasis (Young et al., 2000; Guillod-Maximin et al., 2004; Marty et al., 2005). Our present report provides evidence that astrocytes in the NST respond to changes in glucose availability. Following exposure to conditions of low glucose, a subset of NST-astrocytes and neurons exhibit an increase in intracellular calcium signaling, this is reversed following restoration of normal glucose availability. Both the peak and magnitude of the astrocytic response varied across different levels of low glucose availability. The size of the neuronal response across the different glucose concentrations did not vary significantly. In comparison to adjacent
neurons, astrocytes displayed larger peak responses and response magnitudes, with a reduced latency to respond, following exposure to 0.5 mM glucose concentrations. Note that the activation of astrocytes by reduced glucose was not a smooth linear function; increased activation occurred abruptly at 0.5 mM glucose. This behavior may be a reflection of the operation of an emergency low threshold detection mechanism of the sort driving counter-regulation in response to hypoglycemia (Song et al., 2001; Marty et al., 2007). Furthermore, this astrocytic response to low glucose challenge was independent of neuronal activation, as it was preserved during bath application of TTX. Note that bath application of TTX nearly eliminated the neuronal response to the same low glucose challenge. This suggests that low glucose-mediated increases in neuronal cytoplasmic calcium are produced as a consequence of action potential discharge (Gobel and Helmchen, 2007). In addition, a subset of NST-astrocytes and neurons exhibited an increase in intracellular calcium signaling in response to bath application of the glucoprivic compound 2-DG (5 mM). The onset of this response to glucoprivation was more rapid in astrocytes as compared to adjacent neurons. However, the latencies we observed for astrocytes and neurons are similar to those reported previously for glucose sensitive cells in the hypothalamus or hindbrain (Song et al., 2001; Kang et al., 2004; Balfour et al., 2006). These observations suggest that NST-astrocytes are important sensors of low glucose levels, and that the chemodetection of glucose availability by NST-astrocytes may play an important role in the autonomic regulation of glucose homeostasis.

Astrocyte chemodetection is emerging as an important means by which the brain mediates autonomic homeostasis. Astrocytes in the ventral medulla increase cytoplasmic calcium in response to elevated pCO2. This increase in glial calcium triggers a gliotransmission mechanism that, in this instance, causes the release of ATP onto adjacent respiratory rhythmogenic neurons (Gourine et al., 2010; Marina et al., 2013). Our own work has shown that astrocytes in the NST are activated by proteinase activated receptor (PAR) agonists and, in turn, release glutamate onto adjacent NST neurons (Hermann et al., 2009). In addition to glucoregulatory functions, the NST also suppresses gastric motility through vago-vagal reflex connections with parasympathetic motor neurons (Rogers and Hermann, 2012a). This mechanism may explain in part why bleeding trauma and burn injuries (both conditions
generate large amounts of PAR-activating thrombin) are associated with profound gastrointestinal stasis (Hermann et al., 2009). It remains to be determined whether similar glial-neuronal interactions are present between glucosensitive astrocytes and NST neurons.

In addition, the mechanism by which a reduction in glucose availability signals an increase in cytoplasmic calcium in the NST-astrocyte also remains indeterminate. While astrocytic expression of GLUT2 appears necessary for proper glucose counter-regulation (Marty et al., 2003), the transduction mechanisms often coupled to GLUT2 produce changes in calcium that run in the opposite direction to that observed for astrocytes. For example, pancreatic beta cells respond with an increase in intracellular calcium levels in response to increasing extracellular glucose. Here, glucose is transported through the GLUT2 transporter and metabolized via glycolysis and the tricarboxylic acid cycle to CO$_2$, producing ATP. Increased cellular ATP energy charge acts to phosphorylate and inactivate K$_{ATP}$ channels, leading to depolarization, activation of voltage gated calcium channels and ending in excitation—secretion coupling (Schuit et al., 2001; Holsbeeks et al., 2004; Levin et al., 2004).

However, others have also reported that glucose restriction in cultured astrocytes causes an increase in astrocytic cytoplasmic calcium signals (Arnold, 2005). Similar to our present observations, this increase in cytoplasmic calcium is reversed with the restoration of normal levels of glucose in the perfusion bath. There is evidence suggesting that astrocytes are highly dependent on glycolysis for ATP production; removal of glucose or blocking glucose utilization may rapidly starve astrocytes of glucose for ATP production (Kahlert and Reiser, 2000). With impaired glycolysis and subsequent low ATP following low extracellular glucose availability, the calcium-ATPase pump in the endoplasmic reticulum (ER) of astrocytes fails and ER calcium is released to the cytoplasm (Kahlert and Reiser, 2000; Arnold, 2005). While these results are consistent with our observations in the slice preparation, it is not clear if this mechanism is actually responsible for the cytoplasmic calcium signal seen by our laboratory in in situ NST-astrocytes. For example, the response time for astrocytes in our NST slice preparation is about five-fold faster than that reported for cultured hippocampal astrocytes.

One possibility for the acceleration of this mechanism relative to cultured astrocytes would be the expression of GLUT2 and glucokinase (GK) by glucosensitive NST-astrocytes. Glial expression of both GLUT2 and GK has been reported previously (Garcia et al., 2003; Arluison et al., 2004; Young and McKenzie, 2004; Millan et al., 2010), as well as the specific expression of GLUT2 by a subset of NST-astrocytes (Leloup et al., 1994). Co-expression of these proteins would permit cytoplasmic glucose to flow backward through the GLUT2 transporter following a reduction in extracellular glucose below intracellular glucose concentrations, thus speeding the elimination of fuel for glycolysis, and therefore accelerating ER calcium release. Further studies utilizing single cell qPCR could be employed to directly investigate whether GLUT2 or GK is preferentially expressed by glucosensitive NST-astrocytes as compared to non-glucosresponsive NST astrocytes.

Another possibility which could account for the more rapid astrocytic cytoplasmic calcium signals in our preparation is that NST-astrocytes may contain a much more active low-glucose detection mechanism via direct signaling of low glucose availability by GLUT2. GLUT2 is the only mammalian glucose transporter known to act as a “transceptor,” i.e., a protein which functions as both a transporter and receptor. The receptor function of mammalian GLUT2 is mediated by the large intracellular loop between the 6th and 7th transmembrane domains (Guillemin et al., 2000). Transgenic mice generated to knock down a GLUT2-intracellular loop domain yielded animals that demonstrated an inability to detect glucose but left the GLUT2-dependent glucose transport unaffected (Stolarczyk et al., 2007, 2010). Depression of glucose detection in these animals resulted in increased daily food intake due to increased meal sizes.

The transporter function of GLUT2 has also been linked to the activation of hepatic transcription factor, a carbohydrate response-element binding protein (Uyeda and Repa, 2006). However, it seems unlikely that similar GLUT2-mediated transcriptional regulation in NST-astrocytes could bring about the large and rapid changes in intracellular calcium signaling we observed in response to low glucose availability. Instead, it seems more likely that there may be a direct interaction between the intracellular loop domain of GLUT2 and signaling pathways present in NST-astrocytes; although, at present, there is no information available as to how it might function.

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REFERENCES

Adachi, A., Kobashi, M., and Funahashi, M. (1995). Glucose-responsive neurons in the brainstem. Obes. Res. 3(Suppl. 5), 735S–740S. doi: 10.1002/j.1550-8528.1995.tb00493.x

Andrew, S. F., Dinh, T. T., and Ritter, S. (2007). Localized glucoprivation of hindbrain sites elicits corticosterone and glucagon secretion. Am. J. Physiol. Regul. Integr. Comp. Physiol. 292, R1792–1798. doi: 10.1152/ajpregu.00777.2006

Arluison, M., Guignon, M., Nguyen, P., Thorens, B., Leloup, C., and Penicaud, L. (2004). Distribution and anatomical localization of the glucose transporter 2 (GLUT2) in the adult rat brain—an immunohistochemical study. J. Chem. Neuroanat. 28, 117–136. doi: 10.1016/j.jchemneu.2004.05.009

Arnold, S. (2005). Estrogen suppresses the impact of glucose deprivation on astrocytic calcium levels and signaling independently of the nuclear estrogen receptor. Neurobiol. Dis. 20, 82–92. doi: 10.1016/j.nbd.2005.02.002

Balfour, R. H., Hansen, A. M., and Trapp, S. (2006). Neuronal responses to transient hypoglycaemia in the dorsal vagal complex of the rat brainstem. J. Physiol. 570, 469–484. doi: 10.1113/jphysiol.2005.098822

Bernard, C. (1855). Lecons de Physiologie Experimentale Applique a la Medicine. Paris: Bailliere.

Biag, J., Huang, X., Gou, L., Hintiryan, H., Askarinam, A., Hahn, J. D., et al. (2012). Cyto- and chemoarchitecture of the hypothalamic paraventricular nucleus in the C57BL/6j male mouse: a study of immunostaining and multiple fluorescent trac tract tracing. J. Comp. Neurol. 520, 6–33. doi: 10.1002/cne.22698

Blessing, W. W. (1997). "Anatomy of the lower brainstem," in The Lower Brainstem and Bodily Homeostasis, (Oxford: Oxford University Press), 29–100.

Briski, K. P., and Marshall, E. S. (2000). Caudal brainstem Fos expression is restricted to periventricular catecholamine neuron-containing loci following...
intraventricular administration of 2-deoxy-D-glucose. *Exp. Brain Res.* 133, 547–551. doi: 10.1007/s002210000448

Bucelin, R., and Thorens, B. (2001). Evidence that extrapancreatic GLUT2-dependent glucose sensors control glucagon secretion. *Diabetes* 50, 1282–1289. doi: 10.2337/diabetes.50.12.1282

Cottrell, G. T., and Ferguson, A. V. (2004). Sensory circumventricular organs: central roles in integrated autonomic regulation. *Regul. Pept.* 117, 11–23. doi: 10.1016/j.regpep.2003.09.004

Dallaporta, M., Himmi, T., Perrin, J., and Orsini, J. C. (1999). Solitary tract nucleus sensitivity to moderate changes in glucose level. *Neuroreport* 10, 2657–2660. doi: 10.1097/00001756-199908200-00040

Dodd, G. T., Williams, S. R., and Luckman, S. M. (2010). Functional magnetic resonance imaging and c-Fos mapping in rats following a glucoprivic dose of 2-deoxy-D-glucose. *J. Neurochem.* 113, 1123–1132. doi: 10.1111/j.1471-4159.2010.06671.x

Garcia, M., Millan, C., Balmaceda-Aguilera, C., Castro, T., Pastor, P., Montecinos, H., et al. (2003). Hypothalamic ependymal-glial cells express the glucose transporter GLUT2, a protein involved in glucose sensing. *J. Neurochem.* 86, 709–724. doi: 10.1046/j.1097-4542.2003.01892.x

Geering, J. C., Shin, W., Chimenti, P. C., and Loewy, A. D. (2010). Paraventricular hypothalamic nucleus: axonal projections to the brainstem. *J. Comp. Neurol.* 518, 1460–1499. doi: 10.1002/cne.22283

Gobel, W., and Helmchen, F. (2007). *In vivo* calcium imaging of neural network function. *Physiology (Bethesda)*. 22, 358–365. doi: 10.1152/physiol.00322.2007

Gourine, A. V., Kasymov, V., Marina, N., Tang, F., Figueiredo, M. F., Lane, S., et al. (2013). Astrocyte control through pH-dependent release of ATP. *Science* 329, 571–575. doi: 10.1126/science.1190721

Grill, H. J., and Hayes, M. R. (2012). Hindbrain neurons as an essential hub in the neuroanatomically distributed control of energy balance. *Cell Metab.* 16, 296–306. doi: 10.1016/j.cmet.2012.06.015

Guillod-Maximin, E., Lorsignol, A., Alquier, T., and Penicaud, L. (2004). Acute intracarotid glucose injection towards the brain induces specific c-fos activation in hypothalamic nuclei: involvement of astrocytes in cerebral glucose-sensing in rat nodose ganglia. *J. Physiol.* 588, 617–632. doi: 10.1113/jphysiol.2009.182147

Hermann, G. E., and Rogers, R. C. (2009). TNF activates astrocytes and catecholaminergic neurons in the solitary tract via AMPA receptors: evidence of an atypical neural-glial interaction in the brainstem. *J. Neurosci.* 31, 14037–14045. doi: 10.1523/JNEUROSCI.2855-11.2011

Hermann, G. E., Van Meter, M. J., Rood, J. C., and Rogers, R. C. (2009). Proteinase-activated receptor 2 (GLUT2) and astrocyte-dependent glucose sensors. *J. Clin. Invest.* 115, 3545–3553. doi: 10.1172/JCI26309

McDougal, D. H., Hermann, G. E., and Rogers, R. C. (2011). Vagal afferent stimulation activates astrocytes in the nucleus of the solitary tract via AMPA receptors: evidence of an atypical neural-glial interaction in the brainstem. *J. Neurosci.* 31, 14027–14036. doi: 10.1523/JNEUROSCI.0762-11.2011

Marty, N., Dallaporta, M., Foretz, M., Emery, M., Tarusso, D., Bady, I., et al. (2005). Regulation of glucose secretion by glucose transporter type 2 (GLUT2) and astrocyte-dependent glucose sensors. *J. Clin. Invest.* 115, 3545–3553. doi: 10.1172/JCI26309

Marty, N., Dallaporta, M., and Thorens, B. (2007). Brain glucose sensing, counterregulation, and energy homeostasis. *Physiology (Bethesda)*. 22, 241–251. doi: 10.1152/physiol.00010.2007

McDougall, D. H., Hermann, G. E., and Rogers, R. C. (2011). Vagal afferent stimulation activates astrocytes in the nucleus of the solitary tract via AMPA receptors: evidence of an atypical neural-glial interaction in the brainstem. *J. Neurosci.* 31, 14037–14045. doi: 10.1523/JNEUROSCI.2855-11.2011

Mizuno, Y., and Oomura, Y. (1984). Glucose responding neurons in the nucleus tractus solitarius of the rat: in vitro study. *Brain Res.* 307, 109–116. doi: 10.1016/0006-8993(84)90466-9

Niijima, A. (1984). The effect of D-glucose on the firing rate of glucose-sensitive vagal afferents in the liver in comparison with the effect of 2-deoxy-D-glucose. *J. Auton. Nerv. Syst.* 12, 255–260. doi: 10.1016/0169-2063(84)90221-3

Nimmerjahn, A., Kirchhoff, F., Kerr, J. N., and Helmchen, F. (2004). Sulforhodamine 101 as a specific marker of astroglia in the neocortex in vivo. *Nat. Methods* 1, 31–37. doi: 10.1038/nmeth706

Nickoloff, C., Steinberg, E., Spindel, M. S., and Gripple, F. M. (2006). Sodium-coupled glucose cotransporters contribute to hypothalamic glucose sensing. *Diabetes* 55, 3381–3386. doi: 10.23736/db06-0531

Ritter, S., Dinh, T. T., and Zhang, Y. (2000). Localization of hindbrain-glucoreceptive sites controlling food intake and blood glucose. *Brain Res.* 856, 37–47. doi: 10.1016/S0006-8993(99)02327-6

Ritter, S., Llewellyn-Smith, I., and Dinh, T. T. (1998). Subgroups of hindbrain catecholamine neurons are selectively activated by 2-deoxy-D-glucose induced metabolic challenge. *Brain Res.* 805, 41–54. doi: 10.1016/S0006-8993(98)00655-6

Rogers, R. C., and Hermann, G. E. (2012a). “Brainstem control of gastric function,” in *Physiology of the Gastrointestinal Tract, 4th Edn.*, ed L. R. Johnson (San Diego, CA: Elsevier Academic Press), 861–892. doi: 10.1016/B978-0-12-382026-6.00031-2

Rogers, R. C., and Hermann, G. E. (2012b). Tumor necrosis factor activation of vagal afferent terminal calcium is blocked by cannabinoids. *J. Neurosci.* 32, 5237–5241. doi: 10.1523/JNEUROSCI.6220-11.2012

Rogers, R. C., Nasse, J. S., and Hermann, G. E. (2006a). Live-cell imaging methods for the study of vagal afferents within the nucleus of the solitary tract. *J. Neurosci.* 26, 15070–15078. doi: 10.1523/JNEUROSCI.3949-06.2005

Rogers, R. C., Van Meter, M. J., and Hermann, G. E. (2006b). Tumor necrosis factor potentiates central vagal afferent signaling by modulating ryadnone channels. *J. Neurosci.* 26, 12642–12646. doi: 10.1523/JNEUROSCI.0330-06.2006

Sanders, N. M., and Ritter, S. (2000). Repeated 2-deoxy-D-glucose-induced glucoprivation attenuates Fos expression and glucoregulatory responses during subsequent glucoprivation. *Diabetes* 49, 1865–1874. doi: 10.2337/diabetes.49.11.1865

Schuit, F. C., Huygens, P., Heimberg, H., and Pipeleers, D. G. (2001). Glucose sensing in pancreatic beta-cells: a model for the study of other glucose-regulated cells in gut, pancreas, and hypothalamus. *Diabetes* 50, 1–11. doi: 10.2337/diabetes.50.1.1
Silver, I. A., and Erecinska, M. (1994). Extracellular glucose concentration in mammalian brain: continuous monitoring of changes during increased neuronal activity and upon limitation in oxygen supply in normo-, hypo-, and hyperglycemic animals. *J. Neurosci.* 14, 5068–5076.

Song, Z., Levin, B. E., McArdle, J. J., Bakhos, N., and Routh, V. H. (2001). Convergence of pre- and postsynaptic influences on glucosensing neurons in the ventromedial hypothalamic nucleus. *Diabetes* 50, 2673–2681. doi: 10.2337/diabetes.50.12.2673

Stolarczyk, E., Guissard, C., Michau, A., Even, P. C., Grosfeld, A., Serradas, P., et al. (2010). Detection of extracellular glucose by GLUT2 contributes to hypothalamic control of food intake. *Am. J. Physiol. Endocrinol. Metab.* 298, E1076–E1087. doi: 10.1152/ajpendo.00737.2009

Stolarczyk, E., Le Gall, M., Even, P., Houllier, A., Serradas, P., Brot-Laroche, E., et al. (2007). Loss of sugar detection by GLUT2 affects glucose homeostasis in mice. *PLoS ONE* 2:e1288. doi: 10.1371/journal.pone.0001288

Uyeda, K., and Repa, J. J. (2006). Carbohydrate response element binding protein, ChREBP, a transcription factor coupling hepatic glucose utilization and lipid synthesis. *Cell Metab.* 4, 107–110. doi: 10.1016/j.cmet.2006.06.008

Watts, A. G., and Donovan, C. M. (2010). Sweet talk in the brain: glucosensing, neural networks, and hypoglycemic counterregulation. *Front. Neuroendocrinol.* 31:32–43. doi: 10.1016/j.yfrne.2009.10.006

Yettefti, K., Orsini, J. C., El Ouazzani, T., Himmi, T., Boyer, A., and Perrin, J. (1995). Sensitivity of nucleus tractus solitarius neurons to induced moderate hyperglycemia, with special reference to catecholaminergic regions. *J. Auton. Nerv. Syst.* 51, 191–197. doi: 10.1016/0165-1888(94)00130-C

Yettefti, K., Orsini, J. C., and Perrin, J. (1997). Characteristics of glycemia-sensitive neurons in the nucleus tractus solitarii: possible involvement in nutritional regulation. *Physiol. Behav.* 61, 93–100. doi: 10.1016/S0031-9384(96)00358-7

Young, J. K., Baker, J. H., and Montes, M. I. (2000). The brain response to 2-deoxy glucose is blocked by a glial drug. *Pharmacol. Biochem. Behav.* 67, 233–239. doi: 10.1016/S0091-3057(00)00315-4

Young, J. K., and McKenzie, J. C. (2004). GLUT2 immunoreactivity in gomori-positive astrocytes of the hypothalamus. *J. Histochem. Cytochem.* 52, 1519–1524. doi: 10.1369/jhc.4A6375.2004

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