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ORIGINAL ARTICLE



Changes in neuronal activity across the mouse ventromedial nucleus of the hypothalamus in response to low glucose: **Evaluation using an extracellular multi-electrode array** approach

Lydia Hanna^{1,2} | Tristan J. Kawalek² | Craig Beall² | Kate L. J. Ellacott²

¹Reading School of Pharmacy, University of Reading, Reading, UK

²Institute of Biomedical & Clinical Sciences, University of Exeter Medical School, Exeter, UК

Correspondence

Kate L. J. Ellacott, Neuroendocrine Research Group, University of Exeter Medical School, Level 4 RILD, Barrack Road, Exeter EX2 5DW UK Email: k.ellacott@exeter.ac.uk

Present address

Lydia Hanna, Department of Biological Sciences, Centre for Biomedical Sciences, Royal Holloway University of London, Egham, UK

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Abstract

The hypothalamic ventromedial nucleus (VMN) is involved in maintaining systemic glucose homeostasis. Neurophysiological studies in rodent brain slices have identified populations of VMN glucose-sensing neurones: glucose-excited (GE) neurones, cells which increased their firing rate in response to increases in glucose concentration, and glucose-inhibited (GI) neurones, which show a reduced firing frequency in response to increasing glucose concentrations. To date, most slice electrophysiological studies characterising VMN glucose-sensing neurones in rodents have utilised the patch clamp technique. Multi-electrode arrays (MEAs) are a state-of-the-art electrophysiological tool enabling the electrical activity of many cells to be recorded across multiple electrode sites (channels) simultaneously. We used a perforated MEA (pMEA) system to evaluate electrical activity changes across the dorsal-ventral extent of the mouse VMN region in response to alterations in glucose concentration. Because intrinsic (ie, direct postsynaptic sensing) and extrinsic (ie, presynaptically modulated) glucosensation were not discriminated, we use the terminology 'GE/presynaptically excited by an increase (PER)' and 'GI/presynaptically excited by a decrease (PED)' in the present study to describe responsiveness to changes in extracellular glucose across the mouse VMN. We observed that 15%-60% of channels were GE/PER, whereas 2%-7% were GI/PED channels. Within the dorsomedial portion of the VMN (DM-VMN), significantly more channels were GE/PER compared to the ventrolateral portion of the VMN (VL-VMN). However, GE/PER channels within the VL-VMN showed a significantly higher basal firing rate in 2.5 mmol I⁻¹ glucose than DM-VMN GE/PER channels. No significant difference in the distribution of GI/PED channels was observed between the VMN subregions. The results of the present study demonstrate the utility of the pMEA approach for evaluating glucose responsivity across the mouse VMN. pMEA studies could be used to refine our understanding of other neuroendocrine systems by examining population level changes in electrical activity across brain

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nuclei, thus providing key functional neuroanatomical information to complement and inform the design of single-cell neurophysiological studies.

KEYWORDS

electrophysiology, glucose-sensing, hypothalamus, multi-electrode array, ventromedial nucleus

1 | INTRODUCTION

The brain is critical for maintaining systemic glucose homeostasis, in part by direct sensing of changes in glucose levels. Neurophysiological studies in rodent brain slices have identified glucose-sensing neurones that change their firing frequency in response to alterations in local glucose levels. Glucose-sensing neurones are broadly subdivided into two groups: glucose-excited (GE) and glucose-inhibited (GI). GE neurones show enhanced firing activity in response to higher ambient glucose concentrations, whereas the reverse is true for GI neurones. The hypothalamus is a key brain region mediating glucose homeostasis and glucose-sensing neurones have been identified in several hypothalamic nuclei: the arcuate nucleus (ARC), lateral hypothalamus, paraventricular nucleus and ventromedial nucleus (VMN).¹ Microdialysis studies in rats have shown that glucose concentrations in the ventromedial region of the hypothalamus (VMH, which comprises the ARC and VMN) are approximately 20%-30% that of peripheral euglycaemic ('normal') levels: approximately 1.0-2.5 mmol L⁻¹ (VMH) and 5.5 mmol L⁻¹ (blood). Within the VMH, these can reach 5 mmol L^{-1} during hyperglycaemia and 0.2 mmol L^{-1} during hypoglycaemia.²⁻⁴ Therefore, the glucose concentrations and time courses used in the present study model pathophysiologically relevant variations seen in people with diabetes.

The VMN is one of the best studied hypothalamic nuclei with respect to glucose-sensing neurones, with both GE and GI neurones observed.^{1,5} VMN glucose-sensing neurones play an important role in detecting and reacting to glucose deficit, regulating both the counter regulatory response to hypoglycaemia and glucoprivic feeding.⁶ GE and GI neurones are defined as being directly/intrinsically responsive to alterations in extracellular glucose levels.⁷ Indeed, in the presence of the voltage-gated sodium channel blocker, tetrodotoxin (TTX), direct postsynaptic changes (ie, in the resting membrane potential) are seen in both GE and GI neurones in various brain areas.^{8,9} Three subtypes of non-intrinsic, glucose-sensing VMN neurones have also been described, which are modulated by presynaptic glutamatergic inputs. These extrinsically glucose-sensing neurones include PED neurones, which are presynaptically excited by a decrease (PED) in extracellular glucose levels (from 2.5 to 0.1 mmol L⁻¹ glucose), and PER and PIR neurones, which are either presynaptically excited (PER) or inhibited (PIR) by an increase in extracellular glucose levels (from 2.5 to 5-10 mmol L⁻¹ glucose), respectively.¹⁰

Unlike other hypothalamic nuclei where the neuropeptide/neurotransmitter phenotype of glucose-sensing neurones has already been identified,¹ this is less clear for the VMN. To date, most of the ex vivo slice electrophysiological studies characterising VMN glucose-sensing neurones in rodents have utilised the patch clamp technique, which enables real-time detailed interrogation of the electrical properties of individual glucose-sensing neurones. However, because only a single cell can be studied at a given time, and given that intrinsic GE and GI neurones are estimated to together comprise only approximately 20% of the VMN neuronal population,¹¹ this makes the patch clamp technique both time-consuming and of relatively low yield. Furthermore, because few glucose-sensing neurones can be recorded per brain slice, the investigator, who, in this circumstance, is ultimately searching for specialised glucose-sensing neurones, is potentially introducing unintentional anatomical selection bias in their recordings by targeting parts of the VMN known to be enriched in these cells.

Multi-electrode arrays (MEAs) comprise an electrophysiological tool enabling the electrical activity of large neuronal groups to be simultaneously recorded across multiple electrode sites (channels). This powerful neurophysiological extracellular recording technique is used both in vivo and ex vivo.¹² An advantage of this method compared to others is the high spatial and temporal resolution that it offers, enabling the functional investigation and mapping of complex and heterogeneous nuclei, such as the VMN.^{13,14} Each MEA electrode site can record the electrical activity from a population of neurones: referred to as multi-unit activity (MUA). Use of conventional spike sorting methods allows the activity of single cells (single-unit activity [SUA]) to be discriminated from the MUA.¹⁵ We have previously employed perforated MEA (pMEA) technology for ex vivo hypothalamic recordings from mouse brain slices ¹⁶ because of the improved slice perfusion rate, long-term recording stability and high signal-to-noise ratio.¹⁷ In the present study, we have used an equivalent pMEA extracellular recording approach in ex vivo adult mouse brain slices aiming to objectively evaluate changes in neuronal activity across the dorsal-ventral extent of the medial portion of the VMN in response to changes in extracellular glucose concentration. To our knowledge, this is the first study utilising the MEA system to evaluate glucose responsivity across the mouse VMN neural network.

2 | MATERIALS AND METHODS

2.1 | Animals

Male 10-13 week-old C57BL/6J mice bred at the University of Exeter were used for these studies (n = 5, 1-2 VMN-containing slices per

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animal). Mice were group housed under a 12:12 hour light/dark cycle at 22 \pm 2°C, with unlimited access to standard laboratory rodent diet (EURodent diet [5LF2]; LabDiet; PMI Nutrition International, LLC, Brentwood, MO, USA) and water. Studies were performed in accordance with UK Home Office Regulations as defined in the UK Animals (Scientific Procedures) Act 1986 and were approved by the University of Exeter Animal Welfare Ethical Review Board.

2.2 | Reagents

Unless stated otherwise, all reagents were obtained from Sigma-Aldrich (Poole, UK) or Fisher Scientific (Loughborough, UK).

2.3 | Preparation of brain slices

Mice were killed by cervical dislocation followed by decapitation at zeitgeber time (ZT)3 (with lights on defined as ZT0, lights off as ZT12). Brains were rapidly removed and then, along the coronal plane, sectioned using a Leica VT1200 microtome (Leica Microsystems, Wetzlar, Germany) in ice-cold (~4°C) sucrose-based solution composed of (in mmol L⁻¹): 189 sucrose, 10 D-glucose, 26 NaHCO₃, 3 KCl, 5 MgSO₄·7H₂0, 0.1 CaCl₂ and 1.25 NaH₂PO₄, bubbled with 95% O₂/5% CO₂ mixture. Slices (thickness 300 µm) containing the medial portion of the VMN were collected, starting from bregma –1.22 mm up to –2.18 mm ¹⁸ (for more details, see Anatomical locations section below). Slices containing the medial-VMN (one or two per animal) were used for subsequent recordings. Slices were transferred into a Petri dish containing artificial cerebrospinal fluid (aCSF) composed of (in mmol L⁻¹): 10 D-glucose, 124 NaCl, 3 KCl, 24 NaHCO₃, 1.25 NaH₂PO₄, 1 MgSO₄·7H₂0 and 2 CaCl₂, continuously bubbled with carbogen. After a 20-minute rest at room temperature (~22°C), slices were then transferred into a holding chamber containing oxygenated aCSF with physiologically-relevant brain glucose concentrations (in mmol L⁻¹): 2.5 D-glucose, 7.5 D-mannitol, 124 NaCl, 3 KCl, 24 NaHCO₃, 1.25 NaH2PO₄, 1 MgSO₄·7H₂O and 2 CaCl₂: slices were gradually warmed to ~34°C in the holding chamber and allowed to rest for at least 1 hour.

2.4 | In vitro multi-electrode recordings

MEA recordings from acute brain slices were performed as described previously.^{16,19} After 1 hour of rest. a VMN-containing mouse brain slice was placed recording side down, onto a 60pMEA100/30iR-Ti-gr perforated multi-electrode array (pMEA; Multi Channel Systems, MCS GmbH, Hanover, Germany). These arrays are comprised of 60 electrodes in a 6 × 10 layout (one is a reference electrode), with each electrode 100 µm apart, and with a diameter of 30 μ m, covering a total area of approximately 707 μ m². Thus, anatomically, a single array can cover the entirety of the mouse VMN.¹⁸ Contrast illumination (both underneath and above the brain slice) was used to ensure accurate placement of the VMN on the array. To verify VMN positioning over the pMEA electrode sites, images were taken using a GXCAM-EYE-5 5MP eyepiece camera (GT Vision, Assington Green, UK) and overlaid. The slice was acclimatised in pre-warmed oxygenated aCSF containing 2.5 mmol L⁻¹ glucose in the pMEA recording chamber for at least 30 minutes before the recordings began.

To ensure recording stability, slices were held in place by negative pressure provided by vacuum-generated suction applied via perforations of the MEA and by a weighted slice harp. The pMEA recording chamber was continuously perfused with pre-warmed ($34 \pm 1^{\circ}$ C)

Slice	Total number of VMN channels	Total number of non- VMN channels	Total number of non- parenchymal channels	Number of DM-VMN channels	Number of VL- VMN channels
1	56	0	3	33	23
2	52	3	4	23	29
3	18	39	2	8	10
4	55	0	4	26	29
5	49	6	4	20	29
6	58	0	1	29	29
7	36	18	5	19	17
8	41	16	2	22	19
Total	365	82	25	180	185
Mean per slice	46	10	3	23	23
SEM	5	5	<1	3	3
Median per slice	51	5	4	23	26
IQR	40	<1	2	20	19

TABLE 1 Distribution of recording channels (multi-unit) across the ventromedial nucleus (VMN) of the mouse hypothalamus

Note: Data rounded to the nearest integer. n = 5 mice (one or two slices per mouse). Abbreviations: DM, dorsomedial; IQR, interquartile range; VL, ventrolateral.

 TABLE 2
 Distribution of total identified single-units across the
ventromedial nucleus (VMN) of the mouse hypothalamus

Slice	Total number of single-units	Total number of VMN single-units
1	6	6
2	2	2
3	5	5
4	12	12
5	9	8
6	5	5
7	12	6
8	10	8
Total	61	52
Mean per slice	8	7
SEM	1	1
Median per slice	8	6
IQR	5	5
	-	

Note: Data rounded to the nearest integer where appropriate. n = 5 mice (one or two slices per mouse). Abbreviation: IQR, interquartile range.

oxygenated aCSF (containing 2.5 mmol L⁻¹ glucose) from above and below at a combined rate of 2.5-3 mL min⁻¹. Neural signals were acquired using MC_RACK software (MCS GmbH) as time-stamped action potential waveforms using a MEA1060 system and MEA1060UP amplifier (MCS GmbH). Signals were sampled at 50 kHz, high-pass filtered at 200 Hz (second-order Butterworth) and multi-unit spikes crossing a threshold (usually set at -16.5μ V) were extracted for further analysis.

2.5 Glucose responsiveness

To investigate the glucose-sensing capability of VMN neural networks to changing glucose concentrations, recordings were commenced in 'euglycaemic' 2.5 mmol L⁻¹ D-glucose containing aCSF (referred to as 2.5 mmol L⁻¹ glucose). After a 40-minute baseline recording, slices were then perfused for 15 or 40 minutes in 'hypoglycaemic' 0.1 mmol L⁻¹ D-glucose containing aCSF (referred to as 0.1 mmol L⁻¹ glucose), composed of (in mmol L⁻¹): 124 NaCl, 3 KCl, 24 NaHCO₃, 1.25 NaH₂PO₄, 1 MgSO₄·7H₂O, 0.1 D-glucose, 9.9 D-mannitol and 2 CaCl₂: slices were then allowed to recover in 2.5 mmol L⁻¹ D-glucose containing aCSF. These glucose concentrations were chosen because they reflect VMN glucose concentrations observed during euglycaemia and hypoglycaemia: approximately 1.0-2.5 mmol L⁻¹ and 0.2 mmol L⁻¹, respectively.²

To investigate the glucose-sensing capability of VMN neural networks to a less extreme change in glucose concentration (more in line with an overnight fast 10), after a baseline recording in 2.5 mmol L⁻¹ glucose (as above), slices were then perfused for 15 minutes in 0.7 mmol L⁻¹ D-glucose containing aCSF (referred to as 0.7 mmol L⁻¹ glucose), composed of (in mmol L⁻¹): 124 NaCl, 3 KCl, 24 NaHCO₃,

1.25 NaH₂PO₄, 1 MgSO₄·7H₂0, 0.7 D-glucose, 9.3 D-mannitol and 2 CaCl₂: slices were then allowed to recover in 2.5 mmol L⁻¹ D-glucose containing aCSF.

No synaptic blockers were used during recordings. Although the voltage-gated sodium channel blocker TTX has been useful in patch clamp experiments to distinguish between intrinsic and extrinsic glucosensing capabilities of neurones,^{8,9} we could not use this approach during MEA recordings as bath application of TTX abolishes the extracellularly recorded spontaneous action potentials in VMN-containing brain slices²⁰. However, at the end of a subset of experiments, slices were either treated with bath application of the glutamatergic NMDA receptor agonist, NMDA (20 µmol L⁻¹), to confirm maintained cell responsiveness, and/or were followed by application of TTX (1 μ mol L⁻¹). Given the strength of the spatial coverage of the pMEA, we chose to focus on the responsiveness of glucose-sensing neurones to low glucose across the mouse VMN, rather than on intrinsic/extrinsic glucosensation. For conciseness, glucose-sensing neuronal populations (multi-unit activity) detected by the pMEA electrodes were termed as: 'GE/PER' or 'GI/PED' (the latter includes adapting GI channels described previously by others 21,22).

2.6 Statistical analysis

MUA and isolated SUA (ie, from these multi-unit recordings) were analysed in OFFLINE SORTER, version 3 (Plexon Inc., Dallas, TX, USA) using principal component analysis-based spike sorting, as described previously.¹⁶ Briefly, a distinct cluster of spikes was identified as a single-unit if confirmed by multivariate analysis of variance F statistics and a clear refractory period (>20 ms) in the interspike interval distribution. After spike sorting, the data were then exported to NEUROEXPLORER, version 4 (Nex Technologies, Colorado Springs, CO, USA) and MATLAB R2016b (MathWorks Inc., Natick, MA, USA) for construction of perievent histograms and further analysis. Glucose-responsive units were identified as those in which there was a clear peak or trough during the 0.1 or 0.7 mmol L⁻¹ glucose application that exceeded the upper or lower bounds of the 95% confidence limits from the pre-stimulus spike counts, respectively (10-second bins). All units that maintained a low firing frequency (< 0.2 Hz) throughout the experiment and/or did not respond to alterations in glucose concentration, including during recovery of glucose levels from 0.1 or 0.7 mmol L^{-1} back to 2.5 mmol L^{-1} , were considered non-responsive.

Data in graphs and tables are expressed as the mean ± SE and/or median and interguartile range as appropriate. To compare groups, a paired t test (normally distributed data) or a Wilcoxon signed rank test (non-normally distributed data) was used as appropriate. The Shapiro-Wilk normality test was used to assess whether data were normally distributed. Two-way ANOVA with repeated measures and Bonferroni post-hoc test was used to evaluate the impact of glucose concentration on mean firing frequency between regions of the VMN. Statistical analyses were performed using PRISM (GraphPad Software Inc., San Diego, CA, USA).

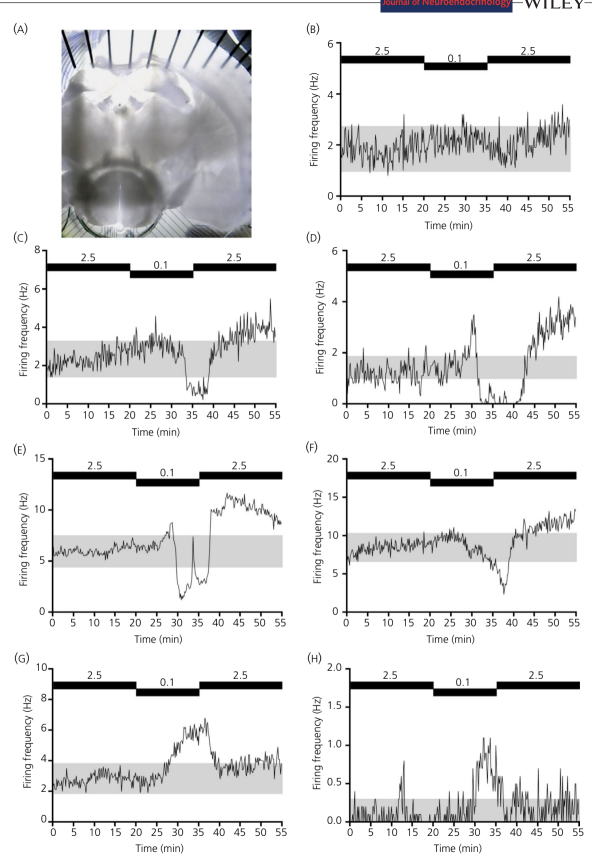


FIGURE 1 Representative examples of multi-electrode array (MEA) captured multi-unit activity across the ventromedial nucleus (VMN)-containing slice. A, Representative image of an acute, coronal-cut mouse VMN-containing slice used for subsequent MEA recording (recording side shown). Representative perievent histograms from non-responsive (B), glucose-excited (glucose-excited/presynaptically excited by an increase) (C-E), late glucose-excited/presynaptically excited by an increase responder (F) and glucose-inhibited (glucose-inhibited/presynaptically excited by a decrease) (G, H) multi-unit channels. Black bars (B-H) indicate bath application of 2.5 and 0.1 mmol L⁻¹ glucose (15 minutes). Grey shading (C-H) represents the 95% confidence limits

2.7 | Anatomical locations

Offline overlay of pMEA electrode sites and respective recorded slice images, with the aid of key landmarks (shape of the hippocampus, third ventricle, median eminence and location of the optic tracts) and reference to the Mouse Brain Atlas,¹⁸ were used to determine the location of each pMEA electrode site/channel: VMN, non-VMN or non-parenchymal (Table 1 [multi-unit] and Table 1 [single-unit] for 2.5 to 0.1 mmol L⁻¹ [40 minutes] studies). Specifically, the relative shape of both the hippocampus and third ventricle was studied to confirm the rostral-caudal location of the mouse VMN-containing slice (from bregma -1.22 mm up to -2.18 mm) and to verify that the cutting plane was not angled. The size and shape of the third ventricle then determined the optimal VMN-containing brain slice(s) for recording, thus ensuring consistency between animals and experiments. Although every effort was made to only include electrodes in the analysis that fell within the VMN, recording locations are only approximate to the VMN and its subdivisions and there is a possibility that some electrode recordings are not from the VMN but, instead, from adjacent regions. Within the mouse hypothalamus, as a result of the challenges associated with accurate delineation of the sub-regions, the VMN was split into a dorsomedial (DM) and ventrolateral (VL) region, with its central region equally split between the two regions. Collectively, this allowed for VMN regional differences to be studied and, importantly, to be distinguished from non-VMN regions, with the latter being excluded from VMN-related analyses (Table 2).

3 | RESULTS

3.1 | Extracellular MEA recordings detect the presence of glucose-sensing channels in the VMN

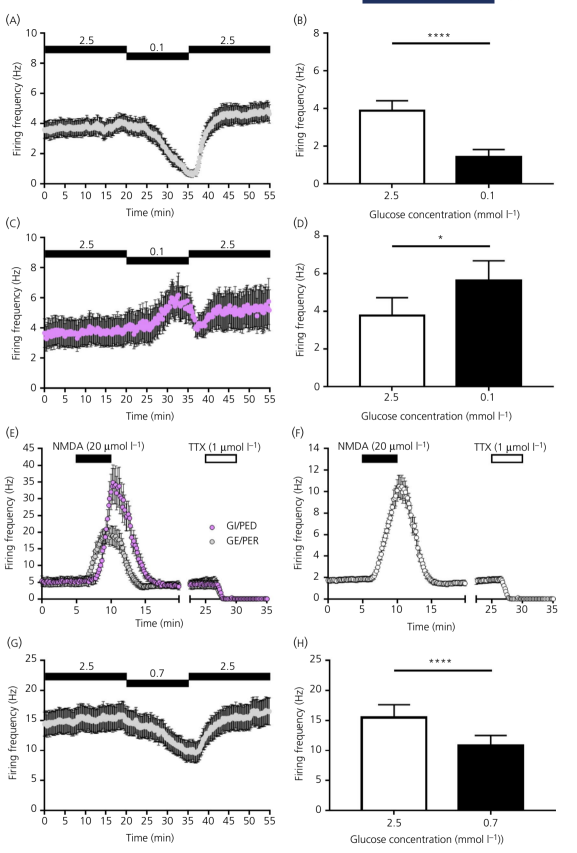
To our knowledge, this is the first study to employ the MEA extracellular recording technique to investigate glucosensation in the VMN.

Using this neurophysiological recording approach, we performed some preliminary experiments to test whether protocols similar to those used in patch clamp experiments could be employed to reveal the presence of glucose-sensing channels in VMN-containing brain slices prepared from mice (Figure 1A). By lowering the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ for 15 minutes, we found that, although most channels did not change their firing frequency (Figure 1B), approximately 15% (n = 31 out of 210 VMN channels: for pooled data, see Figure 2A,B) of VMN channels could be classified as GE/PER (Figure 1C-E). The most commonly observed GE/PER channel response profiles are shown in Figure 1C,D (MUA analysis). Of note, a proportion of VMN channels were not included in any further analysis (~5%; n = 11 out of 210 VMN channels) because they were classified as late GE/PER responders (Figure 1F): despite the clear decline in firing rates during the 0.1 mmol L⁻¹ glucose application, firing rates did not cross the 95% confidence limits (grey shading) within the 15-minute application window; thus, they were not classified as GE/PER channels. Nonetheless, as expected, GE/PER channels (as did late GE/PER responders) displayed a reversible and sustained reduction in firing frequency in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L^{-1} (Figure 2A,B).

Although rarer, we also observed VMN channels that could be classified as GI/PED channels, accounting for < 4% (n = 8 out of 210 VMN channels: for pooled data, see Figure 2C,D) of VMN channels (Figure 1G,H, MUA analysis). By contrast to GE/PER channels, GI/PED channels displayed a reversible increase in firing frequency in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ for 15 minutes. GI/PED channels observed had similar characteristics to non-adapting GI (Figure 1G) and adapting GI profiles (Figure 1H) as described previously²¹; however, we could not (in this or other experiments) stratify the data by the two GI channel subtypes with confidence, as a result of the low channel number.

As a positive control, the ionotropic glutamate (NMDA) receptor agonist NMDA was bath applied to confirm cell/slice responsiveness. All GI/PED (n = 8/8) and 90% of GE/PER (n = 28/31) VMN channels

FIGURE 2 Firing frequency across the ventromedial nucleus (VMN) in response to changing glucose concentrations and to application of NMDA and tetrodotoxin (TTX) (multi-unit analysis). A, Firing frequency over time in response to changing the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ (15 minutes) and back to 2.5 mmol L⁻¹ (indicated by black bars) across all VMN glucose-excited (GE/ presynaptically excited by an increase [PER]) channels. B, Mean firing frequency over the last 5 minutes in 2.5 mmol L¹ glucose vs the last 5 minutes in 0.1 mmol L⁻¹ glucose across all VMN GE/PER channels. ****P < 0.0001 Wilcoxon matched-pairs signed rank test, n = 31 GE/ PER channels (four slices from four mice; total of 210 VMN channels). C, Firing frequency over time in response to changing the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ (15 minutes) and back to 2.5 mmol L⁻¹ (black bars) across all VMN glucose-inhibited (GI/ presynaptically excited by a decrease [PED]) channels. D, Mean firing frequency over the last 5 minutes in 2.5 mmol L¹ glucose vs the last 5 minutes in 0.1 mmol L¹ glucose across all VMN GI/PED channels. *P < 0.05, paired t test, n = 8 GI/PED channels (four slices from four mice; total of 210 VMN channels). E, Mean firing frequency over time across all VMN GE/PER and GI/PED channels in response to bath application (indicated by bars) of the glutamatergic NMDA receptor agonist, NMDA (20 µmol L⁻¹) and the voltage-gated sodium channel blocker, TTX (1 µmol L¹). Multi-unit analysis of n = 28 and n = 8 NMDA responsive VMN GE/PER and GI/PED channels, respectively (four slices from four mice; total of 210 VMN channels). F, Mean firing frequency over time across all NMDA responsive VMN channels (including channels shown in E) in response to bath application (indicated by bars) of NMDA (20 µmol L⁻¹) and TTX (1 µmol L⁻¹). Multi-unit analysis of a total of n = 163 NMDA responsive VMN channels (four slices from four mice; total of 210 VMN channels). G, Firing frequency over time in response to changing the extracellular glucose concentration from 2.5 to 0.7 mmol L^{-1} (15 minutes) and back to 2.5 mmol L^{-1} (indicated by black bars) across all VMN glucose-excited (GE/PER) channels. H, Mean firing frequency over the last 5 minutes in 2.5 mmol L¹ glucose vs the last 5 minutes in 0.7 mmol L¹ glucose across all VMN GE/PER channels. ****P < 0.0001 Wilcoxon matched-pairs signed rank test, n = 27 GE/PER channels (four slices from four mice; total of 215 VMN channels). Data are expressed as the mean ± SEM



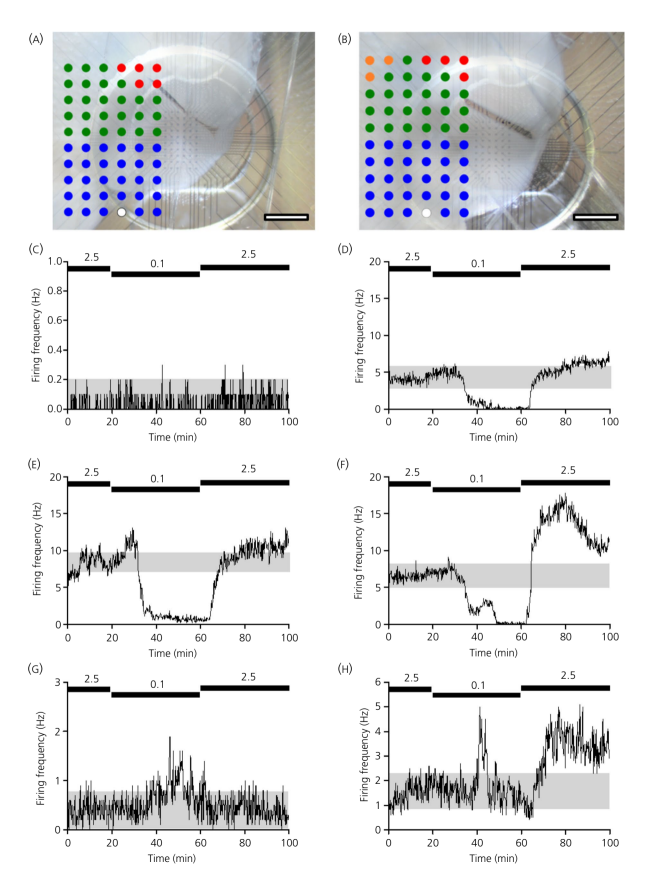
were NMDA responsive (Figure 2E). Indeed, approximately 78% of VMN channels (including those classified as glucose-sensing) were NMDA responsive (n = 163 out of 210 VMN channels) (Figure 2F). The bath application of the voltage-gated sodium channel blocker

TTX completely silenced the recordings (Figure 2E,F), confirming that the acquired signals exclusively reflected sodium-dependent action potentials. This finding also highlights one of the known advantages of the pMEA system: the high signal-to-noise ratio.¹⁷

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We also found evidence of glucose-sensing channels within the mouse VMN in response to less extreme changes in the extracellular glucose concentration: lowering the extracellular glucose concentration from 2.5 to 0.7 mmol L^{-1} for 15 minutes (more closely reflecting brain glucose levels after a 24-hour/overnight fast ¹). With this experimental protocol, we found that, although most channels did not



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slice and multi-unit perievent histograms. A, B, Representative overlaid images of VMN-containing slices, indicating the location of the pMEA electrode sites/channels. Insert shows assigned anatomical location of each channel within the VMN. Green circles = dorsomedial VMN; blue circles = ventrolateral VMN; red circles = non-parenchymal; orange circles = non-VMN; open circle = reference electrode. Scale bars = 700 μ m. Representative perievent histograms from non-responsive (C), glucose-excited (glucose-excited/presynaptically excited by an increase) (D-F) and glucose-inhibited (glucose-inhibited/presynaptically excited by a decrease) (G, H) multi-unit channels. Black bars (C-H) indicate bath application of 2.5 and 0.1 mmol L⁻¹ (40 minutes) glucose. Grey shading (C-H) represents the 95% confidence limits

change their firing frequency, approximately 13% (n = 27 out of 215 VMN channels: for pooled data, see Figure 2G,H) of VMN channels could be classified as GE/PER. These GE/PER channels also displayed a reversible and sustained reduction in firing frequency in response to a lowering the of extracellular glucose concentration from 2.5 to 0.7 mmol L⁻¹ (Figure 2G,H). However, only one channel was classified as an adapting GI/PED channel with this protocol. Furthermore, with this protocol (ie, 2.5 to 0.7 mmol L⁻¹ glucose), there was again a proportion of channels (< 2%; n = 4 out of 215 VMN channels) resembling those previously classified as late GE/PER responders (Figure 1F).

In summary, although we found evidence of glucose-sensing channels within the VMN with an extracellular recording method after a 15-minute application of 0.1 mmol L^{-1} glucose, for the remainder of the studies, an extended 40-minute application was used. This was for two main reasons: (i) to circumvent the issue of omitting late GE/PER (and potentially GI/PED) responders and (ii) to eventually reach a steady-state (ie, plateau) in the response on a population level. We chose the 40-minute timepoint considering that it would likely produce the largest effect across multiple channels and slices, and accepting that some channels may change their frequency more slowly, which we would also capture using this methodology. As such, data discussed from this point onwards is in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L^{-1} for 40 minutes.

3.2 | Most glucose-sensing channels observed in the VMN were glucose-excited (GE/PER)

Using the same pMEA neurophysiological recording approach (Figures 3A,B and 4) combined with the extended (40 minutes) bath application of 0.1 mmol L⁻¹ glucose, approximately 60% of glucose-sensing channels (MUA analysis) detected across the VMN were classified as GE/PER (Figure 3D-F and Table 3), with the most commonly observed GE/PER channel response profiles shown in Figure 3D,E (note the similarities to Figure 1C-E). Although this value is higher than that observed with the previous (15 minutes) experimental protocol (potentially in part as a result of capture and inclusion of late GE/PED responders), these GE/PER channels still displayed a reversible and sustained reduction in firing frequency in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ for 40 minutes (Figure 5).

The mean number of GI/PED channels (MUA analysis) observed across the VMN was < 2% (Figure 3G,H and Table 3): a low percentage similar to that previously observed with the 15-minute 0.1 mmol L^{-1} glucose application (< 4%). GI/PED channels displayed a reversible increase

in firing frequency in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ for 40 minutes. The Gl/ PED channels had characteristics similar to non-adapting Gl (Figure 3G) and adapting Gl profiles (Figure 3H) as described previously.²¹ Rarely, Gl/PED channels showed increased firing upon return to 2.5 mmol L⁻¹ glucose (Figure 3H). MUA analysis revealed a low basal firing rate of Gl/ PED channels, which was increased in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ for 40 minutes (mean \pm SEM: 0.63 \pm 0.25 Hz and 0.79 \pm 0.38 Hz, respectively, n = 6 channels; data not shown). This increase in the mean neuronal firing rate is likely underestimated as a result of the transient responsiveness of adapting Gl channels in low glucose.

3.3 | More glucose-excited (GE/PER) channels were found in the dorsomedial compared to the ventrolateral subdivision of the VMN

Within the VMN, the DM-VMN had significantly more GE/PER channels than the VL-VMN in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L^{-1} for 40 minutes (Figures 6A and 4 and Table 3). There was no significant difference in the distribution of GI/PED channels between these VMN subregions (Figure 4 and Table 3).

3.4 | Glucose-excited (GE/PER) channels in the ventrolateral portion of the VMN displayed a higher spontaneous basal firing frequency

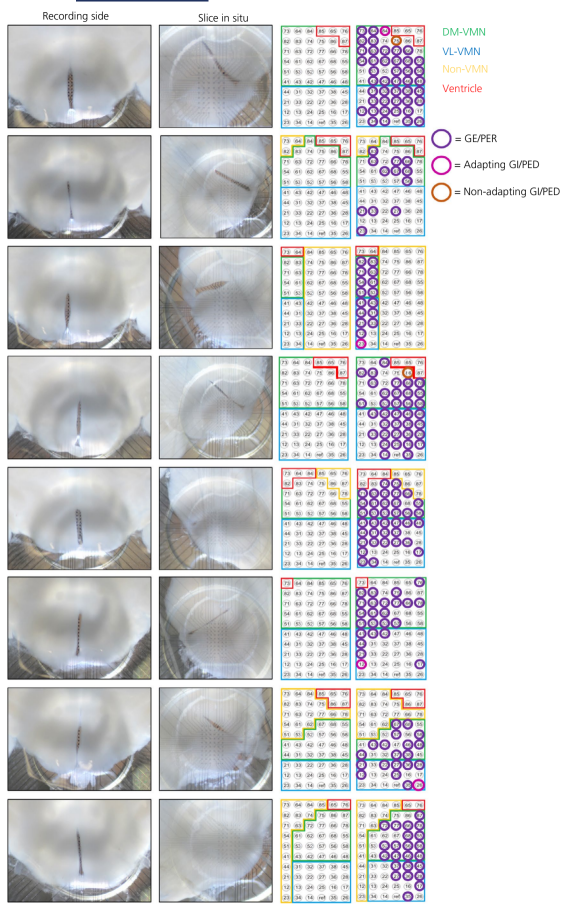
The baseline firing frequency of GE/PER channels (ie, in 2.5 mmol L¹ glucose; MUA analysis) was significantly higher in the VL-VMN compared to the DM-VMN region VMN in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L¹ for 40 minutes (Figure 6B-D). From these data, a representative heat map was generated in which the firing activity across the VMN was plotted relative to anatomical location: this analysis indicated that channels with the highest average basal firing frequency were located in the VL-VMN proximal to the central region (Figure 7).

3.5 | Glucose-inhibited (GI/PED) channels were found across both subdivisions of the VMN

MUA analysis revealed a relatively low number of GI/PED channels (n = 6) in response to a lowering of the extracellular glucose

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FIGURE 4 Multi-electrode array (MEA) channel location and distribution of glucose-sensing channels across all ventromedial nucleus (VMN)-containing slices (40 minutes 0.1 mmol L¹ glucose studies). Slices (thickness 300 μ m) containing the medial portion of the mouse VMN were collected (bregma –1.22 mm up to –2.18 mm) and the recording side of each slice (panels on left) was placed recording side down onto a perforated MEA (pMEA) (see slice in situ panels). Note that the slice is held in place by negative pressure (vacuum-generated suction is gently applied through perforations of the MEA), which may change the appearance of the slice in situ. The numbered grids (panels on the right) indicate location of the pMEA electrode sites/channels: green = dorsomedial (DM) VMN; blue = ventrolateral (VL) VMN; red = non-parenchymal; yellow = non-VMN. The location of the glucose-excited (GE/presynaptically excited by an increase [PER]) channels (multi-unit) on the arrays are indicated by open purple circles. The pink and orange open circles indicate the location of the glucose-inhibited (GI/ presynaptically excited by a decrease [PED]) adapting and non-adapting GI channels, respectively (multi-unit)

TABLE 3 Percentage of glucose-sensing channels (multi-unit), per slice, across the mouse ventromedial nucleus (VMN) of the hypothalamus

Region	Glucose-excited (GE/PER) channels/slice (%, mean ± SEM)	Glucose-inhibited (GI/PED) channels/slice (%, mean ± SEM)	Non-responsive channels/ slice (%, mean ± SEM)	Mean ± SEM channels/slice (n, total channels)
Total VMN	59.49 ± 7.21	1.71 ± 0.66	38.64 ± 7.67	46 ± 5 (365)
DM-VMN	69.07 ± 7.07*	0.86 ± 0.57	30.10 ± 7.07*	23 ± 3 (180)
VL-VMN	51.46 ± 8.78*	2.42 ± 1.33	46.31 ± 9.00*	23 ± 3 (185)

Note: Data from eight slices from five mice, one or two slices per mouse. Number of channels rounded to the nearest integer.

Abbreviations: DM, dorsomedial; VL, ventrolateral; GE, glucose-excited; GI, glucose-inhibited; PER, presynaptically excited by an increase; PED, presynaptically excited by a decrease.

*P < 0.05, Paired t test DM-VMN vs VL-VMN.

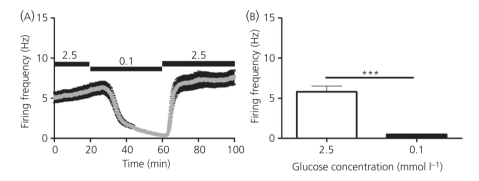


FIGURE 5 Firing frequency across the ventromedial nucleus (VMN) in response to changing glucose concentrations (multi-unit analysis). A, Firing frequency over time in response to changing the extracellular glucose concentration from 2.5 to 0.1 mmol L⁻¹ (40 minutes) and back to 2.5 mmol L⁻¹ (indicated by black bars) across all VMN glucose-excited (GE/presynaptically excited by an increase [PER]) channels. B, Mean firing frequency over the last 5 minutes in 2.5 mmol L⁻¹ glucose vs the last 5 minutes in 0.1 mmol L⁻¹ glucose across all VMN glucose-excited (GE/PER) channels. Data are expressed as the mean ± SEM. ***P < 0.001 Wilcoxon matched-pairs signed rank test, n = 211 GE/PER channels (eight slices from five mice)

concentration from 2.5 to 0.1 mmol L⁻¹ for 40 minutes. GI/PED channels were found in both the DM-VMN and VL-VMN region (Figure 4 and Table 3). However, as a result of the low number of GI/PED VMN channels detected overall, these observations should be interpreted with caution.

3.6 | Analysis of the single-unit data supported the observations of the multi-unit analysis

The SUA supported the MUA data, with 60% of single-units across the VMN being GE/PER (Table 4) and < 7% being GI/PED in response to a lowering of the extracellular glucose concentration from 2.5 to 0.1 mmol L^{-1} for 40 minutes, with the latter being a higher percentage than was seen for the GI/PED MUA analysis. There were insufficient single-units per slice within the VMN sub-regions to provide meaningful analysis of the within VMN distribution.

4 | DISCUSSION

In the present study, we report that, using the pMEA recording technique in ex vivo brain slices, both multi-unit and single-unit analyses confirmed the presence of glucose-sensing neurones across the dorsal-ventral extent of the mouse VMN. This was observed in response to a lowering of the extracellular glucose concentration from 2.5 mmol L^{-1} to either 0.1 or 0.7 mmol L^{-1} glucose

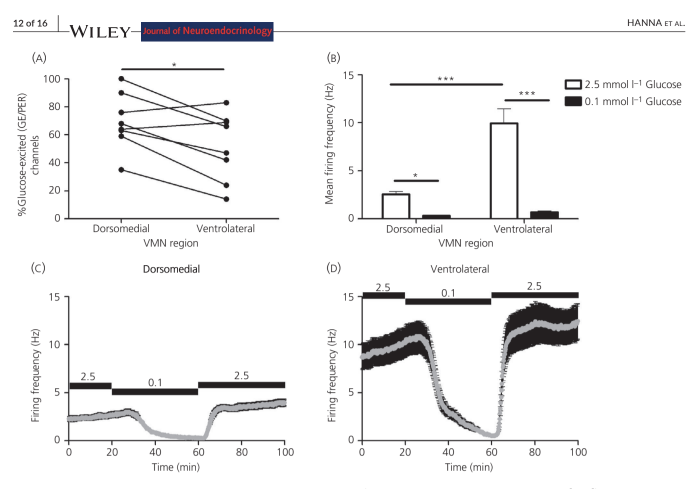


FIGURE 6 Regional differences in firing frequency of glucose-excited (GE/presynaptically excited by an increase [PER]) channels across the ventromedial nucleus (VMN) (multi-unit analysis). A, Percentage of glucose-excited (GE/PER) channels in the dorsomedial compared with the ventrolateral region of the VMN; *P < 0.05, paired t test. B, Mean firing frequency over the last 5 minutes in 2.5 mmol L¹ glucose vs the last 5 minutes in 0.1 mmol L¹ glucose across all VMN GE/PER channels. Mean firing frequency over time in response to changing glucose concentrations from 2.5 to 0.1 mmol L¹ (40 minutes) and back to 2.5 mmol L¹ across GE/PER channels located in (C) the dorsomedial-VMN (DM-VMN) and (D) the ventrolateral-VMN (VL-VMN) region; Two-way repeated measures ANOVA with Bonferroni post-hoc: VMN region, $F_{VMN \text{ region } 1,418} = 32.95$, ***P < 0.001; glucose convcentration, $F_{1,418} = 72.57$, ***P < 0.001; interaction, $F_{1,418} = 26.78$, ***P < 0.001; Bonferroni post-hoc tests, *P < 0.05, ***P < 0.001. Data are expressed as the mean ± SEM. DM-VMN, n = 119, VL-VMN, n = 92 GE/PER channels (eight slices from five mice)

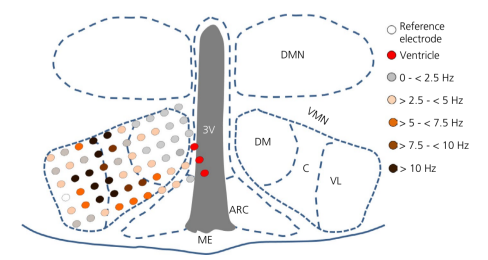


FIGURE 7 Representative heat map of the distribution of glucose-excited (GE/presynaptically excited by an increase [PER]) channels across the mouse ventromedial nucleus (VMN) relative to their respective basal firing frequency in 2.5 mmol L⁻¹ glucose. Glucose-excited (GE/PER) channels within the ventrolateral-VMN (VL-VMN) display a higher basal firing frequency compared to those in the dorsomedial-VMN (DM-VMN): data from 365 channels across eight slices from five mice (multi-unit analysis). ME, median eminence; ARC, arcuate nucleus; 3V, third ventricle; C, central

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TABLE 4 Percentage of glucose-sensing single-units, per slice, across the ventromedial nucleus (VMN) of the hypothalamus

PER)	•		Non-responsive single- units/slice (%, mean ± SEM)	Mean ± SEM single-units/slice (n, total channels)
Total VMN 60.17	7 ± 10.34	6.67 ± 3.27	33.21 ± 10.30	7.1 ± 1.29 (52)

Note: Data from 8 slices from 5 mice, 1-2 slices/mouse. Number of channels rounded to the nearest integer.

Abbreviations: GE, glucose-excited; GI, glucose-inhibited; PER, presynaptically excited by an increase; PED, presynaptically excited by a decrease.

for 15 minutes and from 2.5 to 0.1 mmol L^{-1} glucose for 40 minutes. On this occasion, we focused our main analysis on VMNcontaining slices that were exposed to 0.1 mmol L^{-1} glucose for 40 minutes to enable the detection of the maximal number of GE/ PER units and the achievement of a steady-state in the response to low glucose at the population level.

In agreement with published data utilising single-cell recording techniques in rodent VMN slices, GE/PER channels were much more prevalent than GI/PED channels.^{10,23} At the multi-unit (population) level, although significantly more GE/PER channels were found in the DM-VMN, VL-VMN GE/PER channels showed a significantly higher basal firing frequency (in 2.5 mmol L¹ glucose) than DM-VMN GE/PER channels. MUA and SUA analysis revealed that GI/PED channels are distributed across the subdivisions of the VMN. Further interpretation of the data should be made with caution, as a result of the relatively low number of GI/PED channel subtypes observed in the present study. It is of interest to note the relative quiescence of many of these GI/PED channels (and single-units) during baseline recordings in 2.5 mmol L⁻¹ glucose, in line with findings from patch clamp studies.⁷

The proportion of glucose-sensing channels across the VMN that has been reported previously in single-cell neurophysiological studies in male rodents varies from 50% across the whole VMN ¹⁰ to 20%-80% in the VL-VMN.²¹⁻²³ Data from the present study suggest that, across the whole VMN, the proportion of glucose-sensing channels is 15%-60% depending on the duration of the exposure to 0.1 mmol L⁻¹ glucose, which appears to be an important variable in determining the proportion of glucose-sensing channels. Nonetheless, when single-units were isolated from the multi-unit data for the extended (40 minutes) 0.1 mmol L^{-1} glucose studies, the high percentage of glucose-sensing single-units remains. Some of the reported differences between studies may also be attributed to the age of the animals at the time of study because many studies using single-cell recordings were performed in brain slices from 14-21-day-old juvenile animals, or indeed from species differences because many of the comparable studies have been performed in rats.^{10,23-25} Indeed, separate studies have indicated that there are important species differences with respect to the percentage of glucose-sensing neurones in the VL-VMN: 20% in the juvenile male rat ²³ and 80% in the juvenile male mouse.²¹

The regional differences in number and firing rate of the GE/ PER channels observed in the mouse VMN in the present study may relate to differences in the anatomical location and inputs to the sub-regions within the VMN.²⁶ The higher number of GE/ PER channels in the DM-VMN may be a result of the proximity of this portion of the VMN to the third ventricle. Tanycytes lining the third ventricle have glucose-sensing capacity and extend projections into the VMN,²⁷ enabling direct communication of cerebrospinal fluid glucose levels to DM-VMN neurones. DM-VMN neurones are directly responsive to the key adipostatic hormone leptin via leptin-receptors expressed in this region.²⁸ In addition, the transcription factor steroidogenic-factor 1 (SF-1) is highly expressed in the DM- and central portion of the VMN of the adult mouse,²⁹ and deletion of leptin-receptors from SF-1 neurones in mice using a conditional transgenic knockout approach results in obesity.^{28,30,31} Interestingly, studies in rodents have shown that infusion or microinjection of leptin directly in to the VMH enhances insulin sensitivity and increases glucose uptake in peripheral tissues, including skeletal muscle, heart and interscapular brown adipose tissue ^{32,33}: these effects may be driven by a leptin-activated subset of SF-1 DM-VMN GE neurones.³⁴ In line with these findings, chemogenetic activation of SF-1 VMN neurones stimulates interscapular brown adipose tissue, heart and red-type skeletal muscle glucose uptake.³⁵ Of note, optogenetic modulation of SF-1 VMN neurones also implicates these cells in the regulation of systemic glycaemia, with direct stimulation promoting hyperglycaemia, whereas direct inhibition of these neurones attenuates the counter-regulatory response to insulin-induced hypoglycaemia.³⁶ Collectively, such observations suggest that specific subsets of SF-1 VMN neurones may play distinct roles in systemic glucoregulation. Anatomical tracing studies indicate that the SF-1 VMN neurones critical for the regulation of glycaemia receive inputs from the lateral parabrachial nucleus and mediate their effect on glucose homeostasis via projections to the anterior portion of the bed nucleus of the stria terminalis (aBNST).³⁶ This projection pattern of SF-1 VMN neurones is supported by work conducted by an independent group demonstrating projections from these neurones to sites implicated in regulating sympathetic and autonomic activity,³⁷ including the periaqueductal grey, aBNST, lateral parabrachial nucleus, nucleus of the solitary tract and rostral ventrolateral medulla. Taken together, these findings implicate neurones in the DM-VMN in the central regulation of systemic glycaemia, which may explain the relative abundance of GE neurones in this region of the VMN.

In the present study, the GE/PER channels that showed the highest firing rate in 2.5 mmol L⁻¹ glucose were located in the VL-VMN: the heat map indicates their proximity to the central portion of the VMN. This finding is in agreement with published studies indicating that GE neurones are enriched in the VL-VMN.²³ The central and VL-VMN contain high levels of brain-derived neurotrophic factor, ³⁸ as Y—Journal of Neuroendocrin

well as demonstrating expression of neuroendocrine receptors including oestrogen-receptor α^{39} , progesterone-receptors, ⁴⁰ and rogen receptors⁴¹ and oxytocin receptors.^{42,43} Indeed, although the relative percentage of VL-VMN glucose-sensing neurones is reported to be similar in male and female mice, the proportion of the different subtypes of glucose-sensing neurones in the VL-VMN (GE, adapting GI and non-adapting GI) appears to be sexually dimorphic, which may relate to differential activation of sex steroid hormone receptors impacting neuronal glucose-sensing.^{21,22} Furthermore, the VL-VMN is strongly implicated in mediating sexual behaviour and aggression.⁴⁴ It is not clear why the GE/PER channels in this region showed higher basal firing rates in response to 2.5 mmol L⁻¹ glucose; one possibility could be that, because both sexual behaviour and aggression are energetically expensive, it may make biological sense that a key component in the circuit controlling these behaviours is sensitive to changes in systemic energy availability. Although previous studies have performed detailed electrophysiological characterisation of VMN neurones,⁴⁵ to our knowledge, this is the first report of observed regional differences in basal firing frequency in 2.5 mmol L⁻¹ glucose across the VMN, highlighting a strength of simultaneous recording across the slice, which the MEA approach enables.

The proportion of GI/PED channels detected across the pMEA sites (< 2% MUA, < 7 SUA%; 40 minutes 0.1 mmol L^{-1} glucose) were lower than that reported in previously published studies (single-cell recordings) across the VMN from rats,¹⁰ or the VL-VMN from mice.^{21,22} However, our observations were higher than the 0.4% GI neurones reported in the rat VL-VMN.²³ As such, there appears to be variability in the reported frequency of GI neurones in the VMN. It is possible that, using our extracellular recording method, the prevalence of GI/PED channels is artificially lower than expected because the activity may be masked by the dominating activity of neighbouring GE neurones. This may explain the response profile where we occasionally observed a hump in the firing rate of GE/PER channels during bath application of 0.1 mmol L⁻¹ glucose; for example, during the first 10-15 minutes of low glucose perfusion (Figure 1D at 30 minutes and Figure 3E at 20-30 minutes) or during the nadir phase (Figure 1E at 30-35 minutes and Figure 3F at 40-50 minutes). Nevertheless, in the present study, the percentage of GI/PED channels across the VMN was significantly lower than that of GE/PER channels.

GI neurones in the VMH that specifically express neuronal nitric oxide synthase have been shown to be important for both neuronal glucosensing and regulation of the counter-regulatory response to hypoglycaemia in vivo.⁴⁶ Furthermore, VMN GE and GI neurones have been shown to express SF-1.⁴⁷ A subpopulation of VMN SF-1 neurones has been shown to co-localise with pituitary adenylate cyclase-activating peptide (PACAP) ⁴⁸ and recent work has identified a population of intrinsically GI VMN PACAP-expressing neurones.⁹ Although comprising a relatively small neuronal population, they display a wide distribution across the VMN and are considered to play a role in systemic glucose regulation because chemogenetic stimulation of these neurones in vivo resulted in inhibition of insulin secretion, which led to a reduced glucose tolerance.⁹ Interestingly, GI

VMN PACAP-expressing neurones (15% of which were found to be neuronal nitric oxide synthase-positive) send projections both within the VMH and to other brain areas, including the paraventricular nucleus, lateral hypothalamus, aBNST, paraventricular nucleus of the thalamus and periaqueductal grey,⁹ in line with the findings reported by Meek *et al.*³⁶

In summary, in the present study, we have utilised the pMEA recording technique to provide an unbiased assessment of the percentage and distribution of glucose-sensing neurones across the mouse VMN (ex vivo). Although the pMEA method cannot provide finer, more detailed information regarding the neurophysiological properties of cells, such as can be obtained by the patch clamp technique, it is still a very useful neurophysiological tool for examining both population and single-cell level responses simultaneously across brain nuclei, thus providing, using a high-throughput approach, key functional neuroanatomical information that could complement and inform the design of future single-cell studies. In practical terms, MEA recordings are arguably more user-friendly and technically less demanding to perform than the gold standard patch clamp method and, at the same time, offer a high spatial and temporal resolution.¹⁴ Finally, the in vitro MEA method permits cell-non-invasive, longterm, stable recordings to be performed, and, in combination with optogenetic, chemogenetic, pharmacological and/or calcium imaging approaches, this method can be used to refine our understanding of other neuroendocrine networks.

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CONFLICT OF INTERESTS

The authors declare that they have no conflicts of interest.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

ORCID

Craig Beall Dhttps://orcid.org/0000-0002-4263-0866 Kate L. J. Ellacott https://orcid.org/0000-0001-5261-7465

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