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Single-Cell Imaging of Bioenergetic Responses to Neuronal Excitotoxicity and Oxygen and Glucose Deprivation

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Excitotoxicity is a condition occurring during cerebral ischemia, seizures, and chronic neurodegeneration. It is characterized by overactivation of glutamate receptors, leading to excessive Ca2+/Na+ influx into neurons, energetic stress, and subsequent neuronal injury. We and others have previously investigated neuronal populations to study how bioenergetic parameters determine neuronal injury; however, such experiments are often confounded by population-based heterogeneity and the contribution of effects of non-neuronal cells. Hence, we here characterized bioenergetics during transient excitotoxicity in rat and mouse primary neurons at the single-cell level using fluorescent sensors for intracellular glucose, ATP, and activation of the energy sensor AMP-activated protein kinase (AMPK). We identified ATP depletion and recovery to energetic homeostasis, along with AMPK activation, as surprisingly rapid and plastic responses in two excitotoxic injury paradigms. We observed rapid recovery of neuronal ATP levels also in the absence of extracellular glucose, or when glycolytic ATP production was inhibited, but found mitochondria to be critical for fast and complete energetic recovery. Using an injury model of oxygen and glucose deprivation, we identified a similarly rapid bioenergetics response, yet with incomplete ATP recovery and decreased AMPK activity. Interestingly, excitotoxicity also induced an accumulation of intracellular glucose, providing an additional source of energy during and after excitotoxicity-induced energy depletion. We identified this to originate from extracellular, AMPK-dependent glucose uptake and from intracellular glucose mobilization. Surprisingly, cells recovering their elevated glucose levels faster to baseline survived longer, indicating that the plasticity of neurons to adapt to bioenergetic challenges is a key indicator of neuronal viability.

Key words: bioenergetics; excitotoxicity; single-cell imaging
Likewise, population-based measurements in neuronal cultures often include confounding factors such as contributions from dying neurons or from non-neuronal cells.

To address this deficiency, we used recently developed Förster resonance energy transfer (FRET)-based fluorescent reporters to detect alterations in glucose and ATP concentration, and in AMPK activity at the single-cell level during and after excitotoxicity and oxygen glucose deprivation (OGD)-induced neuronal injury. We show that ATP depletion and recovery during and after excitotoxicity is rapid and shows significant bioenergetic plasticity. We also demonstrate that excitotoxicity resulted in an accumulation of intracellular glucose, and that the ability to rapidly restore intracellular glucose homeostasis after excitotoxicity is a predictor of neuronal survival.

Materials and Methods

Materials. Fetal bovine serum (FBS), minimum essential medium (MEM; plus Earle’s t-glutamine, Invitrogen), Flu/o-4 AM, and tetramethylrhodamine methyl ester (TMRM) were from Invitrogen (BioScience). 5-Amino-imidazole-4-carboxamide [β-D-ribofuranoside (AICAR)] was from Cell Signaling Technology, while Compound C was from Merck Biosciences. Glutamate, NMDA, glycin, MK-801, and all other reagents were obtained from Sigma-Aldrich, unless otherwise stated. The follow-ing media and buffers were prepared: feeding media: cerebellar granule neurons (CGNs; MEM, 32 mm glucose, 18 mm KCl, 10% FBS, and 200 U/ml/200 μg/ml penicillin/streptomycin; cortical neurons (Corticals): MEM, 32 mm glucose, 0.5 mm t-glutamine, 5% FBS, 5% horse serum, and 200 U/ml/200 μg/ml pen/strep; plating media: CGNs: feeding media 2 mm l-glutamine; Corticals: neurobasal medium embryonic, 2% B27, 0.5 mm t-Glutamine, 200 U/ml/200 μg/ml pen/strep; control buffer: 120 mm NaCl, 3.5 mm KCl, 0.4 mm KH2PO4, 5 mm NaHCO3 (Riedel de Haën), 20 mm HEPES, pH 7.4, with 1.5 mm CaCl2, and 15 mm glucose added just before the experiments. Whenever indicated, glucose was either omitted or substituted with 2 mm sodium pyruvate.

Preparation of rat cerebellar granule neurons and mouse cortical neurons. Cerebellar granule neurons were prepared from 7-day-old Sprague-Dawley rat pups of either sex and cortical neurons were prepared from embryonic day 16–18 mouse pups of either sex, as previously described (Dudek et al., 2001; Connolly et al., 2003). Cells were transfected after 5–7 DIV using Lipofectamine 2000 reagent in Opti-MEM Medium (Invitrogen) according to the manufacturer’s instructions. Cells were used 24–48 h after transfection.

Neuronal injury models—excitotoxicity and OGD. Glutamate/glycin or NMDA/glutamine (100/10 μM) was added on stage to CGNs or Corticals, respectively. NMDA receptor activation was ended after 5 or 10 min by the addition of 10 μM MK-801 (Fluka), an NMDA receptor antagonist. Pharmacological compounds [2 μg/ml oligomycin, 12 μM 2-deoxyglucose (2DG), 10 μM compound C (CompC), 100 μM 1,4-dideoxy-1,4-imino-α-arabinitol hydrochloride (DAB), 100 μM 1-deoxyxyritromycin hydrochloride (NIM)] were added to the buffer directly on stage. Excitotoxicity experiments were performed in 15 μM extracellular glucose, unless explicitly stated.

For observation of single-cell kinetics during OGD, cells were placed in a heated stage incubator allowing adjustment of CO2, and O2 concentrations (PeCon). To simulate OGD, media were replaced with ischemic CSF (ICSF; 70 mm NaCl, 70 mm KCl, 1.25 mm NaH2PO4, 2 mm MgSO4, 0.3 mm CaCl, 10 mm sucrose, 5.25 mm NaHCO3, 10 mm TMRM; bubbled with N2), and O2 levels were reduced to 1.5%. After 40–45 min, ICSF media were replaced with conditioned media, and normoxic (20% O2) conditions were restored. During some experiments, focus had to be readjusted after media exchange—out of focus images were not included in the analysis of single-cell kinetics, resulting in gaps in single-cell traces.

Determination of neuronal injury. Following exposure to NMDA, glutamate, or OGD, neurons were allowed to recover in conditioned media at 37°C, 5% CO2 for 24 h before being stained live with Hoechst 33258 stain (1 μg/ml). The extent of neuronal injury compared with sham conditions (control buffer alone) was determined by the percentage of condensed pyknotic nuclei, as described previously (Connannon et al., 2010).

Fluorescence microscopy. The membrane-permeant, cationic fluorescent probe TMRM was used as a measure of changes in the mitochondrial membrane potential. All cells were equilibrated with TMRM (10 nM) in control buffer at least 45 min before imaging (for OGD, cells were maintained in regular media before OGD). For measurements of intracellular calcium, cells were preincubated with Flu/o-4 AM (3 μM) for 45 min. Mineral oil was added on top of the buffer to minimize evaporation.

The ATeam 1.03 FRET-based probe, kindly provided by Dr. Hiroyuki Noji (Osaka University, Osaka, Japan), is a highly specific and sensitive reporter of cytosolic ATP concentration, and is pH insensitive around physiological pH (Imamura et al., 2009). ATP binding to the ε-subunit of the bacterial E/F1-ATP synthase alters the probe conformation, increasing FRET (see Fig. 2A). The FRET-based AMPK activity reporter (AMP-KAR), kindly provided by Dr. Lewis C. Cantley (Harvard Medical School, Boston, MA), contains a synthetic peptide incorporating an AMPK substrate motif. Specific AMPK phosphorylation of the threonine residue on the synthetic peptide alters the conformation of a FRET reporter of intracellular glucose concentration with a linear response range between 0.05 and 9.6 mM (Takahana et al., 2008). Glucose binding to the glucose–galactose-binding protein alters the probe conformation, increasing FRET, as shown in Figure 4A (Tsou et al., 2011). The probe is therefore an indicator of AMPK activity. The FL1+ Pglu700μs68 “glucose-FRET” probe (plasmid #17866, Addgene) is a specific probe of intracellular glucose concentration with a linear response between 0.05 and 9.6 mM (Takahana et al., 2008). Glucose binding to the glucose–galactose-binding protein alters the probe conformation and increases FRET (see Fig. 5A). This plasmid has been optimized to reduce pH sensitivity [enhanced yellow fluorescent protein (YFP) replaced with citrine protein] and other potential sources of artifact (Fehr et al., 2003; Takahana et al., 2008). A control-FRET reporter (FL1+ Pglu- D183A: glucose-FRET probe with a residue-mutation to reduce glucose sensitivity; Takahana et al., 2008), was used as a negative control for glucose sensitivity, and was kindly provided by Dr. Wolf B. Frommer (Carnegie Institute, Stanford, CA). For interexperimental comparison, we normalized all FRET-based glucose measurements (15 and 0 mM extracellular glucose) to an average of the baseline ratiometric fluorescence signal measured in neurons incubated in 0 mM glucose (2 ± 0.6 pyruvate). As further negative controls to exclude potential artifacts affecting the fluorescent proteins, we used a FRET-based plasmid with a caspase–cleavable linker sequence [SCAT3 (enhanced cyan fluorescent protein (eCFP)–DEVD–Venus); Takemoto et al., 2003]. The SCAT3 ratiometric fluorescence signal remained unaltered during excitotoxic
stimulation, which is in line with the reported absence of significant
executioner caspase activity in models of excitotoxicity (Lankiewicz et al.,
2000; Ward et al., 2006).
Glass bottom WillCo-dishes were mounted on a heated stage above a
40× or 63× oil-immersion objective lens. Inverted confocal laser-
scanning microscopes (LSM 510 Meta, Slive duoscan, and 710, Zeiss)
were used with optimized polychromic mirror and filter wheel settings.
Fluorophores were excited using a 405 nm (CFP) or 488 nm diode laser
(YPF, Fluo-4), a 488 nm argon laser (YPF, Fluo-4), a 543 nm helium/
neon laser, or a 561 nm DPSS laser (TMRRM). Emitted fluorescence
was detected through bandpass or long-pass filters for the LSM 510 and Slive
duoscan, according to the maximum emission wavelengths of the fluo-
rophores (Fluo-4, 515 nm; TMRRM, 576 nm; CFP, 475 nm; YPF,
527 nm). In the case of the LSM 710, the spectral detection bands were
selected accordingly. Cells were imaged on stage for at least 10 min before
treatment to measure baseline fluorescence. Images were obtained ap-
proximately every 1 min before, during, and for at least 10 min after
treatment, and at 5 min intervals for longer experiments. Cells were
imaged on stage for up to 16 h after treatment. Efforts were made to
reduce phototoxicity by minimizing exposure time and laser excitation
power.
Image processing. Images were processed using MetaMorph 7.5 64 bit
(Molecular Devices). After background subtraction, YPF channel images
were used to segment transfected cells. The ratiometric values shown in
the traces were obtained by calculating the ratio image of the FRET/
CFP × 1000 for the segmented cells, obtaining the kinetics of the mean
value for each cell. The fluorescence signal referred to as FRET here is the
fluorescent signal emitted in the YPF band after 405 nm excitation. For
fluorescent dyes (TMRRM and Fluo-4 AM), individual cells were isolated
within regions of interest (ROIs), and the average signal intensity within
the ROI was plotted over time. All traces shown are normalized to base-
line (average of the first eight images) and are representative of at least
three independent experiments, unless explicitly stated.
Intracellular glucose measurement by glucose oxidase assay. After 8 DIV,
media in 6-well plates were replaced with control buffer. After 1 h, Cor-
ticals/CGNs were treated with NMDA/glutamate for 3/10 min and
allowed to recover for up to 2 h. Neurons were washed three times in
control buffer (containing no glucose) and lysed in hypotonic lysis buffer
(100 mM Tris, 2.5 mM EDTA, pH 7.75) before being stored at
−80°C. Intracellular glucose levels were measured using the Amplex Red Glucose
Oxidase Assay kit (Invitrogen), following the manufacturer’s instruc-
tions. Glucose oxidase reacts with glucose to produce gluconoacetate
and H2O2. The subsequent reaction of H2O2 with the Amplex Red
Reagent in the presence of horseradish peroxidase generates a red fluo-
rescent oxidation product whose fluorescence emission was measured at
595 nm after excitation at 560 nm.
Statistical analysis. Single-cell data (time-stamp, fluorescence intensity
value) were input to MATLAB R2007B (MathWorks) where trace char-
acteristics (time to depolarization, area under the curve, and duration of
recovery to baseline) were either manually identified or automatically
calculated. In each box-and-whisker plot, the box has the lines at the lower,
median, and upper quartile values. The whiskers extend to the most
extreme values within 1.5 times the interquartile range. Cells lying out-
side this range are indicated by filled circles. Mean ± SEM values are
shown in the bar charts.
Each experiment was performed on neurons from at least three inde-
pendent preparations. To analyze single-cell behavior, each neuron’s re-
sponse was treated as an independent event in statistical analyses.
The occurrence of significant differences between the medians was deter-
dined by Kruskal–Wallis test (followed by a multiple-comparison test)
and rank sum/paired rank sum tests using Bonferroni correction for
multiple pairwise comparisons. Differences in the means were calculated
using the Student’s t test. Pearson and Spearman correlation tests were
performed using the MATLAB function “corr.” A p value of 0.05 was
considered to be significant (*p < 0.05; **p < 0.01). The Kaplan–Meier
curve, which reports the percentage of surviving neurons over time (each
step in the curve represents one depolarization event, lowering the per-
centage), was generated in StataIC 12 (StataCorp). For survival analysis,
ATP depletion during transient excitotoxicity is a rapid,
transient process, and ATP recovery is incomplete after OGD.
We previously demonstrated that the above transient glutamate
exposure depleted ATP levels in CGN populations, with com-
plete ATP recovery within 1–2 h (Ward et al., 2007; Weisová et al.,
2009). To investigate ATP dynamics in our single-cell system, we
transiently transfected neurons with the recently developed
FRET-based reporter of cytosolic ATP levels, ATeam (Imamura
et al., 2009; Fig. 2A). We first confirmed that this tool was capable of
reporting ATP levels in our system, by inhibiting glycolytic and
mitochondrial ATP production (with 2-DG and oligomycin, re-
spectively). As expected, the ATeam fluorescence signal collapsed
only in the presence of both inhibitors (Fig. 2B; we observed
similar behavior for oligomycin addition followed by 2-DG; data
not shown), suggesting that either glycolytic or mitochondrial
ATP production is sufficient to maintain ATP levels in CGNs at
rest. Oligomycin addition in the presence of 0 mM extracellular
glucose (where mitochondrial ATP production is assumed to be
the only form of ATP production) also induced an immediate
collapse of the ATeam fluorescence signal (data not shown).
Surviving neurons from experiments that were performed for <6 h
were removed.
Results
Transient excitotoxic and OGD injury models induce changes in
intracellular calcium and mitochondrial membrane potential
To investigate neuronal NMDA receptor-mediated glutamate ex-
citotoxicity (Castillo et al., 1998; Ward et al., 2000), we tran-
siently exposed rat CGNs (7–9 DIV) to glutamate and its
coagonist glycine (100/10 μM) for 10 min (Fig. 1A), and observed
significantly increased cell death after 24 h, compared with sham
controls (Fig. 1B). We performed time-lapse confocal micros-
copy to monitor single CGNs before, during, and after the exci-
toxic insult. Using Fluo-4 AM, a fluorescent reporter of
intracellular calcium (Ca2+), and TMRRM, a cationic fluorescent
dye reporting changes in mitochondrial membrane potential (ΔΨm), we
confirmed a rapid and transient increase in intracel-
lar Ca2+ and depletion in ΔΨm (Fig. 1C), which is consistent
with previous results (Nicholls et al., 2003; Ward et al., 2007).
This was followed in some neurons by delayed apoptotic death,
characterized by delayed calcium deregulation and depolariza-
tion of ΔΨm (Fig. 1C, dotted trace). A second model of neuronal
NMDA receptor-mediated excitotoxicity, where mouse Corticals
were exposed to NMDA/glycine (100/10 μM) for 5 min (Fig. 1D),
exhibited similar Ca2+ and ΔΨm dynamics (D’Orsi et al., 2012)
and also increased neuronal death after 24 h (Fig. 1E).
In addition, we extended our investigation to OGD, a widely
used in vitro model of ischemic stroke, consisting of both excito-
toxic and non-excitotoxic features (Goldberg and Choi, 1993).
Implementing an oxygen regulation system that allowed us to
continuously monitor single neurons within our confocal mi-
croscopy setting, we found that exposure of cortical neurons to
OGD for 45 min (Fig. 1F) significantly increased cell death after
24 h (Fig. 1G). Similar to the observations in the pure excitotoxic-
ity models above, we found that ΔΨm was transiently depleted
during treatment, and recovered after restoration of normoxic
and normoglycemic conditions (Fig. 1H). Interestingly, and in
contrast to the excitotoxicity models, an initial increase in Ca2+
was quickly re-equilibrated during treatment, while a secondary
increase in Ca2+ was measured after reoxygenation, with a more
gradual return to baseline.
similarly during and after transient NMDA stimulation (Connolly et al., 2012). Single-cell fluorescence intensity traces of CGNs stained with Fluo-4 (black lines, a reporter for intracellular calcium) and TMRM (gray lines, a reporter of mitochondrial membrane potential), illustrative traces of seven cells are representative of at least three independent experiments. Glutamate rapidly and transiently increased intracellular calcium, and mitochondrial membrane potential decreased throughout glutamate exposure, and more slowly returned to baseline. The dotted trace shows a CGN undergoing delayed calcium deregulation and mitochondrial membrane depolarization (rapid loss of TMRM signal). Two cells were removed from the right hand trace for clarity. The thick gray bar indicates the time of OGD.

We next used the ATeam reporter to monitor cytosolic ATP levels in single neurons undergoing transient excitotoxicity. Strikingly, we detected a rapid ATP depletion in CGNs during glutamate stimulation followed by a similarly rapid recovery to baseline (to within 10% of baseline fluorescence). ATP levels subsequently remained constant until depolarization, when ATP levels rapidly and transiently increased intracellular calcium. Mitochondrial membrane potential decreased throughout glutamate exposure, and more slowly returned to baseline. The dotted trace shows a CGN undergoing delayed calcium deregulation and mitochondrial membrane depolarization (rapid loss of TMRM signal). Two cells were removed from the right hand trace for clarity. The thick gray bar indicates the time of OGD.

ATP recovery after transient glutamate exposure involves active mitochondria

Mitochondria are the major source of neuronal ATP production, and their function can be severely compromised by excessive calcium buffering, such as during excitotoxicity (Nicholls and Budd, 2000). Interestingly, however, our data demonstrated that cytosolic ATP levels recovered significantly earlier than the TMRM signal (data not shown). Utilizing TMRM as a reporter of mitochondrial function, this suggested that, despite some form of remnant mitochondrial dysfunction, CGNs were nevertheless able to completely restore cytosolic ATP. To investigate whether mitochondria were capable of participating in this rapid ATP recovery subsequent to glutamate exposure, we preincubated CGNs in glucose-free buffer supplemented with Na pyruvate (2 mM), bypassing glycolysis and providing substrate directly to the mitochondria. Cytosolic ATP levels in these cells were depleted to 10% of baseline fluorescence within 10 min of stimulation termination in both CGNs exposed to glutamate (Fig. 2G) and Corticals exposed to NMDA (Fig. 2H), ATP remained significantly lower than baseline in Corticals even up to 60 min after the termination of OGD (Fig. 2I).

Figure 1. Single-neuron time-lapse imaging experimental models of transient NMDA receptor-mediated excitotoxicity and OGD. A, Transient glutamate excitotoxicity experimental treatment paradigm. After 7–9 DIV, CGNs were mounted on a confocal microscope and exposed to glutamate (100 μM) and its coagonist glycine (10 μM) for 10 min. Neurons were imaged for up to 16 h after treatment. B, Cell death after 24 h (as quantified by pyknotic nuclei counts) was significantly higher in neurons transiently exposed to glutamate (Glut) compared with sham conditions (p = 4 × 10−14). C, Single-cell fluorescence intensity traces of CGNs stained with Fluo-4 (black lines, a reporter for intracellular calcium) and TMRM (gray lines, a reporter of mitochondrial membrane potential), illustrative traces of seven cells are representative of at least three independent experiments. Glutamate rapidly and transiently increased intracellular calcium. Mitochondrial membrane potential decreased throughout glutamate exposure, and more slowly returned to baseline. The dotted trace shows a CGN undergoing delayed calcium deregulation and mitochondrial membrane depolarization (rapid loss of TMRM signal). Two cells were removed from the right hand trace for clarity. The thick gray bar indicates the time of glutamate exposure. D, Transient NMDA-mediated excitotoxicity experimental treatment paradigm. A model similar to that of CGNs exposed to glutamate, 7–9 DIV Corticals were exposed to NMDA (100 μM) and glycine (10 μM) for 5 min. E, Levels of cell death were significantly higher in Corticals exposed to NMDA compared with those exposed to sham conditions (p = 8 × 10−14). Single-cell Fluo-4 and TMRM traces of Corticals exposed to NMDA have been described previously (D’Orsi et al., 2012). F, OGD experimental treatment paradigm. The 7–9 DIV Corticals were mounted on a confocal microscope, and exposed to hypoxic and hypoglycaemic conditions for 45 min, as described in the Materials and Methods section. G, Levels of cell death were significantly higher in neurons exposed to OGD compared with those exposed to sham conditions (p = 4 × 10−14). H, Single-cell fluorescence intensity traces of Corticals stained with Fluo-4 (black lines) and TMRM (gray lines). The onset of OGD induced a transient increase in intracellular calcium levels and mitochondrial membrane depolarization. Restoration of normoxic/normoglycemic conditions induced a second transient increase in intracellular calcium, and a recovery of mitochondrial membrane potential. Illustrative traces of seven cells are representative of three independent experiments. The thick gray bar indicates the time of OGD.

Figure 2. ATP is depleted but rapidly recovers in two experimental models of transient excitotoxicity and this recovery is incomplete after OGD. A, B, The ATeam FRET-based reporter used in single-cell experiments is a reporter of intracellular ATP concentration. A, Binding of ATP to the $\nu$-subunit of the bacterial F$_0$/F$_1$-ATP synthase linker protein increases the fluorescence ratio of FRET/CFP [illustration adapted from Imamura et al., 2009 (permission to publish received from authors)]. B, Control experiments demonstrated the complete collapse of ATeam ratiometric fluorescence, to $\sim$40% of baseline, only after the inhibition of both glycolytic (2-DG) and mitochondrial ATP production (oligomycin), confirming the suitability of the ATeam probe for tracking intracellular ATP levels in our system. C, Single-cell fluorescence ratio and intensity traces of CGNs transfected with the ATeam FRET-based reporter of ATP concentration (black lines), stained with TMRM (gray lines), and exposed to glutamate for 10 min (thick gray bar). Illustrative traces of five cells are representative of five independent experiments. Depolarization was closely accompanied by a rapid decrease of ATP concentration. Tracked neurons were classified as nonsurvivors (dotted traces) if they depolarized within 6 h, and survivors if they remained viable beyond this time (Fig. 6). D, Fluorescence and differential interference contrast (DIC) images of a representative CGN, transiently transfected with the ATeam FRET-based reporter, before, during, and after glutamate exposure, and after mitochondrial membrane depolarization. Color scale represents ratiometric fluorescence values normalized to baseline. E, Single-cell fluorescence ratio and intensity traces of Cortical neurons transfected with the ATeam FRET-based reporter of ATP concentration (black lines), stained with TMRM (gray lines) and exposed to NMDA for 5 min (thick gray bar). Illustrative traces of nine cells are representative of three independent experiments. F, Single-cell fluorescence ratio and intensity traces of Corticals transfected with the ATeam FRET-based reporter of ATP concentration (black lines), stained with TMRM (gray lines) and exposed to OGD for 45 min (thick gray bar). Illustrative traces of six cells are representative of four independent experiments. G–I, Intracellular ATP concentration before, during, and 10/60 min after exposure to glutamate (29 cells from five experiments; G), NMDA (13 cells from three experiments; H), and OGD (20 cells from four experiments; I). G, H, ATP was significantly depleted during glutamate and NMDA exposure ($p = 4 \times 10^{-4}$ and $p = 1 \times 10^{-4}$, respectively) and recovered within 10 min. I, ATP was significantly depleted during OGD ($p = 2 \times 10^{-7}$) but did not completely recover within 60 min after treatment ($p = 1 \times 10^{-5}$ and $p = 6 \times 10^{-5}$ for 10 and 60 min time points, respectively).
To further investigate the role of mitochondrial ATP production in this process, we treated CGNs in 15 mM glucose with oligomycin (2 μg/ml), an inhibitor of ATP synthase and therefore of mitochondrial ATP production, 60 min before glutamate exposure. Oligomycin had no measurable effect on ATP concentration in neurons at rest \( (p = 2 \times 10^{-5}) \), confirming that glycolysis is sufficient to maintain ATP levels in these circumstances \( (Budd and Nicholls, 1996; Vergun et al., 2003; Jekabsons and Nicholls, 1996) \).

Figure 3. Mitochondrial ATP production is required to rapidly restore ATP to baseline levels after excitotoxic stress. A, B, Removal of glucose from the buffer (with supplementation of 2 mM pyruvate (Pyr)) did not prevent the rapid postglutamate recovery of cytosolic ATP. A, ATeam fluorescence ratio and TMRM average fluorescence intensity traces of CGNs transfected with the ATeam probe, incubated in glucose-free buffer supplemented with pyruvate and exposed to glutamate for 10 min. Illustrative traces of four cells representative of three independent experiments. Subsequent addition of oligomycin completely depleted ATP levels. B, ATP levels of CGNs in glucose-free media supplemented with 2 mM pyruvate were significantly depleted during glutamate exposure \( (17 \text{ cells from three experiments}; p = 2 \times 10^{-5}) \) but recovered within 10 min, similar to CGNs in 15 mM glucose. CGNs were also treated with 2-DG to competitively inhibit glycolytic ATP production. ATP levels in these CGNs recovered but remained significantly depleted 10 min after glutamate exposure \( (23 \text{ cells from three experiments}; p = 2 \times 10^{-5} \text{ and } p = 4 \times 10^{-4} \text{ before compared with subsequent time points}) \). Time points on the x-axes refer to glutamate treatment. C–F, Oligomycin, an inhibitor of ATP synthase, did not induce observable changes in resting ATP concentration, but significantly exacerbated subsequent glutamate-induced ATP depletion, and inhibited the duration and extent of post-gluatamate recovery. C, ATeam fluorescence signal and TMRM average fluorescence intensity traces of CGNs treated with oligomycin \( (10 \mu \text{M}, \text{solid black bar}) \) and subsequently exposed to glutamate for 10 min \( (\text{thick gray bar}) \). Illustrative traces of eight cells are representative of three independent experiments. D, ATP levels in oligomycin-treated CGNs before, during, and 10 and 60 min after glutamate exposure. Oligomycin significantly exacerbated ATP depletion during glutamate exposure and inhibited the rapid recovery \( (31 \text{ cells from three experiments}; p = 9 \times 10^{-11}, 8 \times 10^{-11}, 5 \times 10^{-11} \text{ comparing before with subsequent time points}) \). E, F, Some oligomycin-treated CGNs were capable of partially restoring ATP levels after glutamate exposure. In these cells, however, ATP recovered significantly slower \( (p = 2 \times 10^{-5}; \text{E}) \) and to a lesser extent \( (p = 2 \times 10^{-2}; \text{F}) \) than cells exposed to glutamate alone \( (\text{Control}) \). The extent of ATP recovery was also affected by 2-DG \( (p = 4 \times 10^{-5}) \), although not to the same extent as with oligomycin \( (p = 3 \times 10^{-3}) \) comparing 2-DG with oligomycin.)
AMPK activity in single neurons is rapidly and transiently increased during excitotoxicity, and is decreased during OGD

AMPK is involved in the maintenance of bioenergetic homeostasis (Hardie et al., 2012) and, hence, may regulate energetic recovery after an excitotoxic or OGD-induced energetic crisis. We and others have previously demonstrated that AMPK activity rapidly and transiently increases protein levels of active, threonine-172-phosphorylated AMPK during glutamate excitotoxicity, indicating higher AMPK activity (McCullough et al., 2005; Weisová et al., 2009; Concannon et al., 2010; Davila et al., 2012). To directly monitor AMPK activity in our single-cell system, we used a FRET-based reporter of AMPK activity. The probe incorporates a synthetic peptide that can be phosphorylated by AMPK, altering the probe conformation and increasing FRET, and therefore the ratiometric fluorescent signal (Tsou et al., 2011; Fig. 4A). As a positive control for this probe, we confirmed that the administration of the AMP “mimetic” AICAR or exposure of 2-DG to CGNs and Corticals, respectively, increased the AMPKAR ratiometric fluorescence signal, consistent with the AMPK elevation after these drug exposures observed by other measurements (Fig. 4B). Using this probe, we then found that AMPK activity rapidly and transiently increased during glutamate treatment in CGNs (Fig. 4C,D) and during NMDA treatment in Corticals (Fig. 4E). Notably, in some neurons, AMPK activity began to decrease before termination of the excitotoxic stimulus, suggesting that AMPK activity was no longer required in these cells. In contrast, AMPK activity levels were slightly decreased during OGD treatment in Corticals (Fig. 4F). In all treatment paradigms, AMPK activity rapidly and completely returned to baseline within 10 min after stimulus termination (Fig. 4G–I).

Intracellular glucose increases after excitotoxicity

The rapid recovery of ATP levels indicated that neurons were well capable of satisfying the metabolic requirements for survival in the acute phase after transient excitotoxic insults. However, sources of energy are needed to cope with the excitotoxic injury, and to restore the ionic and metabolic balance. Indeed, previous data from our group demonstrated increased glucose uptake during excitotoxic injury in CGNs in a population-based biochemical assay (Ward et al., 2007), and suggested a role for AMPK-induced GLUT-3 surface expression in mediating this effect (Weisová et al., 2009). We therefore wanted to investigate intracellular glucose levels in our single-cell system, and transiently transfected neurons with a FRET-based reporter of intracellular glucose concentration, glucose-FRET (Takanaga et al., 2008; Fig. 5A). To confirm the sensitivity of the reporter to glucose, we increased the extracellular glucose concentration and detected a corresponding stepwise increase in the emitted glucose-FRET ratiometric fluorescence signal (Fig. 5B).

We initially anticipated that glucose consumption via glycolysis would be increased during excitotoxicity to address the ATP depletion, and that this demand would outweigh any potential elevation in glucose uptake. However, in contrast to this expected decrease in intracellular glucose, we consistently observed a glutamate-mediated increase in CGNs (Fig. 5C–E). As an additional control experiment, we investigated responses of neurons transfected with a mutated version of the glucose-FRET reporter, where one residue of the glucose–galactose binding protein had been mutated to reduce the sensitivity of the reporter to glucose (termed control-FRET herein; Takanaga et al., 2008). We confirmed that the control-FRET reporter was insensitive to changes in extracellular glucose at the levels used in our system (15 mM; data not shown). Control-FRET-transfected CGNs exposed to glutamate for 10 min exhibited a slight increase in the emitted ratiometric fluorescence during the glutamate exposure period only, which may be due to glutamate-induced cellular alterations other than glucose, but otherwise showed no response (Fig. 5F).

To further corroborate the finding of a glutamate-induced increase in intracellular glucose during and after glutamate exposure, we measured intracellular glucose in neuronal populations using an assay based on the conversion of glucose to a fluorometric product via the glucose oxidase enzyme. This assay confirmed that glucose was indeed increased during and shortly after glutamate stimulation in CGNs incubated in 15 mM glucose (Fig. 5G).

We also detected an elevation of intracellular glucose in single Corticals after exposure to NMDA, with kinetics similar to CGNs exposed to glutamate (Fig. 5H–I). Interestingly, OGD did not significantly affect the glucose-FRET fluorescence signal (Fig. 5J,K). Together, these results indicated that abundant intracellular glucose was available to restore ATP after excitotoxic insults, and that substrate depletion was not rate limiting for ATP production in our system.

The excitotoxicity-induced increase in intracellular glucose originates from extracellular, AMPK-dependent glucose uptake and intracellular glucose mobilization

We previously linked glutamate-mediated glucose uptake to elevated AMPK activity (Weisová et al., 2009). Our single-cell data here indeed confirmed a concomitance of transient intracellular glucose, elevated AMPK activity in both excitotoxicity models, and, in turn, demonstrated that no significant glucose elevation was observed during OGD, where AMPK activity was decreased. Hence, to further investigate the role of AMPK activity in glucose levels during excitotoxicity, we treated CGNs with CompC (10 μM), an ATP-competitive inhibitor of AMPK. The initial administration of CompC decreased the glucose-FRET signal, confirming that AMPK regulated intracellular glucose concentration in neurons at rest (Fig. 6A). Of note, the glutamate-induced increase in the glucose-FRET signal was reduced in the presence of CompC (compare Figs. 6B, 5E), yet the signal nevertheless remained elevated beyond stimulus termination (Fig. 6B). This
Figure 4. AMPK activity increases in single neurons during transient glutamate- and NMDA-mediated excitotoxicity, and decreases during OGD. A, B, The AMPKAR FRET-based reporter used in single-cell experiments is a reporter of AMPK activity. A, Phosphorylation of the synthetic peptide by AMPK increases the fluorescence ratio of FRET/CFP, indicating an increase in AMPK activity (illustration adapted from Tsou et al., 2011 [license to publish from Elsevier (#3415451067680)]). B, Exposure of CGNs to 2.5 mM AICAR (31 cells from three experiments) or cortical neurons to 12 mM 2-DG (17 cells from three experiments) increased the fluorescence ratio of the AMPKAR reporter, indicating its suitability for the monitoring of AMPK activity levels in CGNs and Corticals (*p < 10^{-10} and **p < 10^{-9}, respectively). C, D, Glutamate induced a transient increase in AMPK activity. C, Single-cell ratiometric fluorescence signal and intensity traces of CGNs transfected with AMPKAR (black lines), stained with TMRM (gray lines) and exposed to glutamate for 10 min (thick gray bar). Illustrative traces of six cells are representative of four independent experiments. D, Fluorescence and differential interference contrast (DIC) images of a representative CGN, transiently transfected with AMPKAR, before, during, and after glutamate exposure, and after mitochondrial membrane depolarization. Color scale represents ratiometric fluorescence values normalized to baseline. E, NMDA also induced a transient increase in AMPK activity. Single-cell fluorescence ratio and intensity traces of CGNs transfected with AMPKAR (black lines), stained with TMRM (gray lines) and exposed to NMDA for 5 min (thick gray bar). Illustrative traces of nine cells are representative of four independent experiments. F, AMPK activity decreased during OGD. Single-cell fluorescence intensity traces of CGNs transfected with AMPKAR (black lines), stained with TMRM (gray lines), and exposed to OGD for 45 min (thick gray bar). Illustrative traces of eight cells representative of two independent experiments. G–I, Box-plots of AMPK activity before, during and 10 min after exposure to glutamate (31 cells from four experiments; G), NMDA (20 cells from four experiments; H), and OGD (11 cells from two experiments; I). G, H, AMPK activity was significantly elevated during glutamate and NMDA exposure (*p = 6 × 10^{-16} and **p = 1 × 10^{-15}, respectively) and recovered within 10 min (**p = 1 × 10^{-15} and **p = 2 × 10^{-15}, respectively). I, In contrast, AMPK activity was transiently decreased during OGD (p = 1 × 10^{-7}) but again returned to baseline within 10 min (p = 2 × 10^{-10}).
Figure 5. Intracellular glucose is increased in single neurons after transient NMDA receptor-mediated excitotoxicity. A. The glucose-FRET-based reporter used in single-cell experiments is a reporter of intracellular glucose concentration. A, Binding of glucose to the glucose–galactose binding protein increases the fluorescence ratio of FRET/CFP (Takanaga et al., 2008). B, Increasing the extracellular glucose concentration increased the emitted fluorescence ratio in cortical neurons, indicating the suitability of the glucose-FRET probe to report intracellular glucose levels. C, Glutamate induced a transient increase in intracellular glucose. C, Fluorescence intensity traces of individual CGNs transfected with the glucose-FRET reporter (Figure legend continues.)
suggested that, while the glutamate-mediated increase in intracellular glucose was partially AMPK mediated, there nevertheless remained an AMPK-independent fraction. We next wanted to investigate whether the excitotoxicity-induced increase in intracellular glucose was entirely driven by extracellular sources and therefore incubated CGNs in glucose-free media supplemented with pyruvate. We indeed measured a reduced increase in the glucose-FRET signal, confirming that the observed glucose increase was primarily coming from extracellular sources (compare Figs. 6C, 5E). Interestingly, however, the glucose-FRET signal still remained elevated, even beyond stimulus termination, in the absence of extracellular glucose, pointing to an additional and intracellular glucose source for disposal during excitotoxic conditions. Notably, as measured by TMRM signal, these cells showed a recovery of the mitochondrial membrane potential to baseline (to an average of 1.2 ± 0.13 after 10 min), which was comparable to that observed in neurons maintained in 15 mM glucose (1.09 ± 0.05 after 10 min), and remained viable for the duration of the experiments. Similar glucose and TMRM responses were also recorded after the removal of both glucose and pyruvate from the extracellular media, giving evidence that any such internal glucose source may be sufficient to maintain mitochondrial function and neuronal viability after excitotoxic injury (data not shown).

Interestingly, a recent study (Saéz et al., 2014) suggested that neurons are able to mobilize glycogen, and that glycogen metabolism was protective in primary neurons exposed to hypoxia. To test this hypothesis, we inhibited glycogen metabolism with two pharmacological compounds, DAB and NJM (inhibitors of glycogen phosphorylase and α-glucosidase, respectively). Indeed, this inhibition of glycogen metabolism reduced the degree and the time for which the glucose-FRET signal remained elevated, comparable to those responses measured with the control-FRET probe (compare Figs. 6D, 5F), suggesting that glycogen metabolism may contribute to the intracellular glucose increase during excitotoxicity.

(Figure legend continued.) (black lines), stained with TMRM (gray lines), and exposed to glutamate for 10 min (thick gray bar). Illustrative traces of eight cells representative of three independent experiments. D, Fluorescence and differential interference contrast (DIC) images of a representative CGN transiently transfected with the glucose-FRET reporter before, during, and after glutamate exposure, and after mitochondrial membrane depolarization. Color scale represents ratiometric fluorescence values normalized to baseline. E, Glucose-FRET signal in CGNs incubated in 15 mM extracellular glucose before, during, and after glutamate exposure. The FRET signal was normalized to an average of the baseline ratiometric fluorescence signal measured in neurons incubated in 0 mM glucose. The glucose-FRET signal was significantly increased during and after glutamate exposure (32 cells from three experiments; p = 4 × 10^{-2}, 2 × 10^{-1}, and 0.02 comparing before to subsequent time points). F, The ratiometric fluorescence signal of the control-FRET reporter, a mutated version of the glucose-FRET reporter with reduced glucose sensitivity, was increased only during glutamate exposure (18 cells from two experiments; p = 1 × 10^{-2}). G, Measurement of intracellular glucose levels at the population level with a glucose-oxidase fluorometric assay confirmed an increase in glucose levels during and after glutamate exposure in CGNs (three independent experiments; p = 0.04, 0.03, 0.02 comparing before to subsequent time points). H, Fluorescence ratio and intensity traces of individual cortical neurons transfected with the FRET-based reporter of intracellular glucose concentration (black lines), stained with TMRM (gray lines) and exposed to NMDA for 5 min (thick gray bar). Illustrative traces of nine cells representative of three independent experiments. I, The glucose-FRET signal was significantly increased during and up to 15 min after NMDA exposure (19 cells from three experiments; p = 1 × 10^{-2}, 1 × 10^{-1}, 1 × 10^{-1}, and 0.03 comparing before to subsequent time points). J, Fluorescence ratio and intensity traces of individual Corticals transfected with the glucose-FRET reporter (black lines), stained with TMRM (gray lines), and exposed to OGD for 45 min (thick gray bar). Illustrative traces of seven cells representative of three independent experiments. K, The glucose-FRET signal was not significantly altered during or after OGD (10 cells from three experiments).

Neurons that show a prolonged intracellular glucose elevation die earlier

We finally were interested whether the acute responses observed during excitotoxicity were related to the duration of neuronal survival after the insult. We therefore distinguished neurons with sustained TMRM signal >6 h after glutamate exposure, designated “survivors,” from those with a loss of TMRM within this time (“nonsurvivors”; Fig. 2C, dotted line traces). Interestingly, the area under the ATP curve, the minimum ATP levels, or the speed of ATP recovery did not significantly differ between survivors and nonsurvivors (data not shown). This suggested that, once neurons recovered from the acute injury phase, the extent of the glutamate-induced ATP depletion was not a critical determinant or a correlate of the duration of subsequent survival. Similarly, the extent of AMPK activity did not differ between survivors and nonsurvivors (data not shown).

In contrast to ATP concentration and AMPK activity, we surprisingly found that glucose levels in nonsurvivors recovered significantly slower than in those neurons that remained viable after 6 h (survivors; Fig. 6E). Indeed, the recovery of glucose levels was significantly slower and showed a higher cell-to-cell heterogeneity than the recovery of ATP concentration and AMPK activity (Fig. 6F). To further investigate this, we calculated the correlation between the duration of glucose recovery, determined as the time from stimulus termination to recovery to baseline; and the survival time, measured from stimulus termination to loss of TMRM signal. We found that a more rapid return of glucose to baseline, but not other parameters such as maximum glucose amount, significantly correlated with prolonged survival (Spearman correlation coefficient = −0.52; p = 0.039). Additionally, neurons that recovered quicker survived longer than those in which recovery lasted longer (Fig. 6G). From these data, we concluded that the plasticity of neurons to rapidly restore glucose homeostasis was a predictor of neuronal survival.

Discussion

The present study is the first to investigate experimentally, from a single-cell perspective, the behavior of ATP and glucose levels and AMPK activity in single primary neurons undergoing transient NMDA receptor-mediated excitotoxicity or OGD. Our results provide the following findings: using fluorescent protein fusion plasmids based on FRET, we demonstrated in two excitotoxicity models that ATP depletion and recovery to energetic homeostasis, along with AMPK activation, were surprisingly rapid responses. We observed rapid recovery of neuronal ATP even in the absence of extracellular glucose or with inhibition of glycolytic ATP production, demonstrating the remarkable plasticity of neurons in excitotoxic conditions. Of note, we also demonstrated that unimpaired mitochondria were critical for this fast and complete energetic recovery. Using an injury model of OGD, we identified a similarly rapid bioenergetic response, yet with incomplete ATP recovery and decreased AMPK activity. Surprisingly, we also measured a glutamate and NMDA-mediated increase in intracellular neuronal glucose, providing a potential source of energy after glutamate exposure. Further single-cell experiments revealed that this glucose increase originated from extracellular, AMPK-dependent glucose uptake and from intracellular glucose sources, again demonstrating the significant plasticity of primary neurons under excitotoxic conditions. Furthermore, we demonstrated that the ability to more quickly restore glucose levels to baseline was indicative of neuronal survival.

Previous population studies demonstrated that neurons exposed to transient excitotoxicity are able to recover ATP levels,
confirming our single-cell data (Ankarcrona et al., 1995; Ward et al., 2007; Weisova et al., 2009). Population measurements, however, can be skewed by an unsynchronized response, and by contributions from glial cells and dying neurons, resulting in apparently slower kinetics (Loewer and Lahav, 2011). Our experiments showed that after recovery, ATP was maintained at a constant level until cells eventually underwent mitochondrial membrane depolarization. Strikingly, ATP levels were recovered, and were subsequently maintained, both in the absence of extracellular glucose and under conditions of glycolysis inhibition, highlighting the ability of neurons to adapt to varying bioenergetic stressors. Indeed, it has been shown that the ATP produc-

Figure 6. Glutamate-induced glucose elevation originates from extracellular AMPK-dependent glucose uptake and from intracellular glucose mobilization, and glucose-FRET recovery is delayed in CGNs that die earlier. A, Single-cell traces of the ratiometric glucose-FRET fluorescence signal in CGNs treated with compound C (10 μM) 60 min before glutamate treatment. Illustrative traces of six cells representative of three independent experiments. B, Compound C depleted intracellular glucose levels in CGNs at rest (p = 0.004), and reduced but did not completely inhibit the subsequent glutamate-induced glucose-FRET ratio increase (11 cells from three experiments, p = 5 × 10^{-3}, 1 × 10^{-2}, 5 × 10^{-2} comparing before to subsequent time points). C, CGNs incubated in 0 mM glucose also increased their glucose-FRET signal during and after glutamate exposure (20 cells from four experiments, p = 5 × 10^{-3}, 4 × 10^{-3}, 5 × 10^{-3} comparing before to subsequent time points), although this increase was substantially reduced compared with CGNs incubated in 15 mM glucose (compare with Fig. 5E). D, Inhibition of glycogen metabolism (with DAB plus NJM, inhibitors of glycogen phosphorylase and α-glucosidase, respectively) markedly reduced the period for which glucose remained elevated after glutamate stimulation (13 cells from two experiments, p = 1 × 10^{-3} and 5 × 10^{-3}, comparing before to subsequent time points). E, The glucose-FRET signal recovered significantly slower in cells that survived >6 h (survivors; p = 0.027). F, The recovery of the glucose-FRET signal took significantly longer than that of ATP (p = 6 × 10^{-1}) and AMPK activity (p = 9 × 10^{-1}), as measured by the ATeam and AMPKAR reporters, respectively. G, Kaplan–Meier curve demonstrating that CGNs that recovered their glucose-FRET signal to baseline in <24 min survived longer than those whose recovery lasted >24 min. The 95% confidence interval bands are shown.
tion machinery can respond quickly to changing conditions (Klingenberg, 2008). Rodriguez-Rodriguez et al. (2012) demonstrated that, after excitotoxicity, glucose is directed away from the pentose phosphate pathway toward ATP production via glycolysis. We demonstrate that, while mitochondrial ATP production was not necessary for survival in CGNs at rest, it nevertheless played an important role in ATP recovery after glutamate excitotoxicity. Although Ca$^{2+}$ sequestration compromises mitochondrial function during excitotoxicity (Nicholls and Budd, 2000), pyruvate supplementation has previously been shown to be protective against excitotoxic injury in CGNs (Vergun et al., 2003; Jekabsons and Nicholls, 2004; but see Khodorov et al., 2012). Our data suggested that mitochondria function sufficiently well to both reduce the initial extent of ATP depletion, and to rapidly and completely restore cytosolic ATP after termination of the excitotoxic stimulus. Indeed, cytosolic Ca$^{2+}$ influx, leading to increased mitochondrial Ca$^{2+}$, has been shown to elevate TCA cycle enzymes under physiological conditions, thereby increasing mitochondrial respiration (Hansford and Castro, 1985; McCormack et al., 1990; Gunter et al., 1994). In excitotoxic conditions, increased respiration (Jekabsons and Nicholls, 2004; Garcia et al., 2005) may also elevate mitochondrial ATP production to address the need for energy, emphasizing that, while mitochondria may be impaired during excitotoxicity, they still maintain an active role in ATP recovery. Increased excitotoxic severity may eventually, however, also induce further mitochondrial dysfunction, increasing the likelihood of necrosis (Kushnareva et al., 2005).

AMPK is activated during energetic stress, and restores energetic homeostasis by increasing ATP production pathways while reducing ATP consumption (Hardie et al., 2012). Intriguingly, AMPK activity increased rapidly during excitotoxic injury, but returned to baseline before stimulus termination in some cells, suggesting that AMPK activity was no longer necessary in these cells. AMPK activity has been shown to be both prosurvival and proapoptotic, depending on cell type and the extent of injury, among other factors (McCullough et al., 2005; Weisová et al., 2009; Concannon et al., 2010; Davila et al., 2012). Here, we did not find any correlation between the acute AMPK response measured during glutamate exposure and the subsequent neuronal survival. In addition, we did not observe prolonged AMPK activity in our system after glutamate exposure. It is therefore possible that the intrinsic neuronal response to AMPK activation, rather than the magnitude of the AMPK activation itself, is a determinant of cell fate. However, we need to note that the relative increase in AMPKAR fluorescence ratio during excitotoxicity was low (~1.2-fold), and that we cannot discount the possibility that AMPK activity may remain chronically elevated at levels below the detection limit achievable with this system. In addition, levels of AMPK activity may differ between the cytosol and other intracellular compartments, such as the nucleus (McCullough et al., 2005; Ju et al., 2011; Kodiha et al., 2011; Davila et al., 2012), and this will be investigated in future work.

We also measured single-cell bioenergetics in neurons undergoing OGD, a model of ischemic neuronal injury that has both excitotoxicity-dependent and excitotoxicity-independent components (Goldberg and Choi, 1993). Of note, our single-cell analysis demonstrated that calcium dynamics differed in our OGD treatment paradigm compared with NMDA receptor-mediated excitotoxicity, with a second increase in intracellular calcium measured after restoration of normoxic and normoglycemic conditions. In contrast to excitotoxic injury, ATP levels also did not completely recover after OGD. This is in agreement with population-based ATP measurements in primary neurons after OGD (Iijima et al., 2003). As mitochondrial ATP production cannot occur in the absence of oxygen, this would largely affect ATP levels in this treatment paradigm. A potential calcium over-load after reperfusion may also impact ATP recovery, necessitating additional ATP consumption to restore ionic homeostasis. Subsequent experiments demonstrated that AMPK activity decreased during OGD and only recovered after OGD termination. This decrease in AMPK activity may also affect the ability of the neuron to restore ATP levels. Hence, despite the fact that OGD-induced injury is sensitive to NMDA receptor antagonists (Goldberg and Choi, 1993), the nonexcitotoxic components induced by the hypoxic and hypoglycemic insult impacts on the ability of neurons to recover their bioenergetics.

Using a FRET-based fluorescent reporter, we measured an accumulation of intracellular glucose in single neurons during NMDA receptor-mediated excitotoxicity. Intracellular glucose concentration is a balance between processes that increase glucose and those that decrease glucose. Decreased glucose consumption in combination with steady or increased glucose import could explain the observed net glucose increase in our system. Curiously, however, the removal of extracellular glucose or the inhibition of AMPK with compound C reduced but failed to completely prevent this increase, suggesting additional intracellular glucose sources during excitotoxicity, arising, for example, from glycogen metabolism. While glycogen is normally not present in neurons, they are capable of the synthesis and lysis of glycogen (Vilchez et al., 2007). Interestingly, glycogen metabolism was recently shown to be protective in primary neurons under conditions of hyponoxia (Saez et al., 2014). Indeed, the inhibition of glycogen metabolism in our system reduced the time for which the glucose-FRET signal remained elevated, suggesting that the breakdown of glycogen may contribute to the extracellular glucose-independent increase in glucose during excitotoxicity.

Finally, we found that glucose recovery to baseline level was significantly slower than that observed for Ca$^{2+}$, ATP, and AMPK, suggesting that excess glucose accumulated in the cell during glutamate stimulation. This accumulation may provide the cell with a “fool-proof” mechanism to address latent ATP demand. Perhaps surprisingly, however, we observed that a slower glucose recovery correlated with shorter survival. It is possible that the delayed glucose recovery directly causes earlier death. Accumulated intracellular glucose has been shown to induce aberrant glycogen synthesis in telencephalic neurons, leading to apoptosis (Vilchez et al., 2007). Likewise, slower recovery to baseline may indicate reduced shuttling of glucose into the pentose-phosphate cycle, which ultimately may decrease the antioxidant defense mechanisms during excitotoxic injury (Rodriguez-Rodriguez et al., 2012). Excessive glucose may itself be toxic by increasing reactive oxygen species production, or may lead to increased lactate production and acidosis. Decreased glucose consumption leading to elevated glucose levels may also be an indicator of downstream mitochondrial defects such as Ca$^{2+}$-induced inhibition of oxidative phosphorylation (Nicholls and Budd, 2000; Kushnareva et al., 2005), although our data indicate that mitochondrial function is retained during the period of glucose elevation. Possibly the most straightforward explanation, however, is that a rapid recovery of glucose homeostasis points to an ability of the neuron to adapt and respond to bioenergetic stressors, and that this plasticity is indicative of underlying neuronal resistance to excitotoxic injury.

In summary, this study is the first to investigate bioenergetic dynamics in single neurons undergoing transient excitotoxicity or oxygen and glucose deprivation. Investigating ATP and glu-
The bioenergetic recovery of excitotoxic neurons was rapid and highly plastic, and that mitochondrial ATP production participated in this recovery. Our data also indicate that the plasticity of neurons to rapidly adapt to bioenergetic challenges may be a key indicator of neuronal viability.

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