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Intracellular ATP Influences Synaptic Plasticity in Area CA1 of Rat Hippocampus via Metabolism to Adenosine and Activity-Dependent Activation of Adenosine A<sub>1</sub> Receptors

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The extent to which brain slices reflect the energetic status of the in vivo brain has been a subject of debate. We addressed this issue to investigate the recovery of energetic parameters and adenine nucleotides in rat hippocampal slices and the influence this has on synaptic transmission and plasticity. We show that, although adenine nucleotide levels recover appreciably within 10 min of incubation, it takes 3 h for a full recovery of the energy charge (to ø0.93) and that incubation of brain slices at 34°C results in a significantly higher ATP/AMP ratio and a threefold lower activity of AMP-activated protein kinase compared with slices incubated at room temperature. Supplementation of artificial CSF with D-ribose and adenine (Rib/Ade) increased the total adenine nucleotide pool of brain slices, which, when corrected for the influence of the dead cut edges, closely approached in vivo values. Rib/Ade did not affect basal synaptic transmission or paired-pulse facilitation but did inhibit long-term potentiation (LTP) induced by tetanic or weak theta-burst stimulation. This decrease in LTP was reversed by strong theta-burst stimulation or antagonizing the inhibitory adenosine A<sub>1</sub> receptor suggesting that the elevated tissue ATP levels had resulted in greater activity-dependent adenosine release during LTP induction. This was confirmed by direct measurement of adenosine release with adenosine biosensors. These observations provide new insight into the recovery of adenine nucleotides after slice preparation, the sources of loss of such compounds in brain slices, the means by which to restore them, and the functional consequences of doing so.

Introduction

The use of brain slices has revolutionized the study of the mammalian CNS, and they have now become a standard preparation in many laboratories and in many areas of neuroscience. Hippocampal brain slices are particularly widely used for studies into the fundamental properties of synaptic transmission and plasticity.

However, it is an unavoidable fact that their preparation is associated with ischemia (decapitation) and tissue trauma (dissection/slice cutting), which will affect metabolic status and result in departure from the in vivo state. Indeed, a substantially compromised energetic state of brain slices at the time of cutting has been demonstrated (Fredholm et al., 1984; Whittingham et al., 1984b), with high energy phosphate levels (ATP, phosphocreatine) in brain slices being as much as 50% lower than their in situ values (Thomas, 1957; Whittingham et al., 1984a; Schurr and Rigor, 1989). Accordingly, basal conditions in hippocampal slices have been described as reflecting a post-ischemic recovery state (Hossmann, 2008). However, as described as far back as the 1950s (McIlwain et al., 1951; McIlwain, 1952), brain slices do show remarkable metabolic recovery after preparation, and it is now common practice to allow a period of incubation (usually 1 h) before they are used for experiments.

Given the widespread use and importance of brain slices to neuroscience, the aims of our study were threefold: first, to assess the metabolic status of brain slices by studying the temperature-dependent recovery and stability of adenine nucleotide levels and energetic parameters, including the activity of AMP-activated protein kinase (AMPK), an enzyme involved in regulation of cellular energy homeostasis and exquisitely sensitive to the cellular ATP/AMP ratio; second, to investigate the potential causes for the reduced ATP content of brain slices compared with reported in vivo values and to evaluate whether the lower ATP levels are attributable to a lack of adenine nucleotide precursors by incubating slices with the free purine base adenine and the sugar precursor of adenylates, D-ribose; finally, to test whether elevated slice ATP levels change the electrophysiological properties of the tissue, such as the probability of transmitter release or the induction and expression of long-term potentiation (LTP).
Our findings provide new insights into the energetic status of brain slices: they show that the loss of ATP precursors is responsible for the decreased ATP content of brain slices and that, by supplementing the artificial CSF (aCSF) with adenine and d-ribose, the recovery of tissue ATP levels can be improved. However, this has measurable consequences in terms of greater activity-dependent release of extracellular adenosine and, via activation of adenosine A1 receptors (A1R), the raising of the threshold for the induction of long-term potentiation.

**Materials and Methods**

**Preparation of brain slices.** Male Sprague Dawley rats (17–27 d old) were killed by cervical dislocation in accordance with Schedule 1 of the United Kingdom Government Animals (Scientific Procedures) Act 1986 and with local ethical review procedures. Sagittal brain slices (400 μm thick), composed of hippocampus and overlying neocortex, were prepared under standardized conditions in ice-cold aCSF containing 10 mM Mg2+ - using a Microm HM 650 V microtome as described previously (Dale et al., 2000; Frenguelli et al., 2007). Slices were either analyzed immediately during cutting for their purine nucleotide content or transferred to the recording chamber or an incubation chamber (50–100 ml) (Edwards et al., 1989) and submerged in continuously circulating, oxygenated standard dCSF at room temperature (22 ± 0.5°C) or 34 ± 1°C. The composition of the standard dCSF solution included the following (in mM): 124 NaCl, 3 KCl, 2 CaCl2, 26 NaHCO3, 1.25 NaH2PO4, 10 d-glucose, and 1 MgSO4·PH 7.4 (with 95% O2/5% CO2).

In a separate set of experiments, nucleotide concentrations of (1) the intact hippocampus and cortex and (2) slices of varying thickness were analyzed. For this purpose, one hemisphere was dissected and separated hippocampus and cortex, and the other hemisphere was used to cut slices of varying thickness (200–3600 μm). Because with the micropotome used no slices >1500 μm could be cut automatically, 3600 μm had to be measured with a guide. For these experiments, all nucleotide extractions were performed immediately after preparation (zur Nedden et al., 2009).

**Nucleotide extraction.** To determine the total adenine nucleotide (TAN) content of 400 μm brain slices, two slices for each time point [from time of cutting (time 0) to 5 h after cutting at 10 min, 30 min, 1 h, 2 h, 3 h, and 5 h] were transferred into ice cold aCSF to stop enzymatic activities. To minimize transfer of aCSF into the reaction mixture, slices were removed with a small spatula into a 1.5 ml microcentrifuge tube containing 1 ml of 5% perchloric acid (PCA). Nucleotide extraction was performed as described in detail previously (zur Nedden et al., 2009). Extracts were neutralized by a threefold organic extraction with 1 ml of tri-n-octylamine dissolved in 1,1,2-trichlorotrifluoroethane (1:1; v/v). The protein pellet was resuspended in 1 ml of 0.5 M NaOH, and the protein concentration was determined by Bradford assay, with bovine serum albumin (BSA) as standard.

For analysis of the TAN content of whole hippocampus and neocortex and 1200, 1500, and 3600 μm slices, the tissue was first homogenized in 500 μl of 5% PCA. The amount of this suspension containing 20 mg wet weight (equivalent to four 200 μm, two 400 μm, or one 800 μm slice) of the tissue was mixed with 5% PCA to a final volume of 1 ml and neutralized as described above.

We have shown previously that snap freezing in liquid N2 and freeze thawing of brain tissue results in a degradation of adenine nucleotides and an underestimation of the energy charge (zur Nedden et al., 2009). For this reason, nucleotides were only extracted from fresh brain tissue and were analyzed on the same day.

**Protein extraction.** For each time point, two to three brain slices were placed in ice-cold aCSF to stop enzymatic activities. Slices were homogenized in 100 μl of protein lysis buffer with a Kontes pellet pestle motor (Sigma-Aldrich). The suspension was centrifuged (30 min, 4°C, 16,000 × g), and the supernatant was stored at ~80°C for kinase assays and Western blot analysis. The composition of the protein lysis buffer was as follows: 50 mM Tris–HCl, pH 7.5, 0.1 mM EDTA, 1 mM EDTA, 1% Triton X-100, 1 μM sodium orthovanadate, 50 mM sodium fluoride, 5 mM sodium pyrophosphate, 270 mM sucrose, 0.1% β-mercaptoethanol, 0.02% sodium azide, and 1 protease inhibitor tablet for 16.6 ml of lysis buffer. HPLC. For analysis of purine nucleotides and nucleosides, an ion pair reversed-phase HPLC method with tetrabutylammonium hydrogen sulfate (TBAHS) was used, as described previously (zur Nedden et al., 2009). Analytical separation was performed on a Supelcosil LC-18-T reversed-phase column (150 × 4.6 mm; inner diameter, 3 μm), with a gradient profile from 100% buffer A (65 mM potassium phosphate, pH 6.0, 4 mM TBAHS) to 100% buffer B (65 mM potassium phosphate, pH 6.0, 25% methanol) in 15 min. Peak identities were confirmed by comparison of the retention times of sample peaks with peaks of standard compounds, spiking the samples with individual standards and by comparison of the UV spectra with standard compounds. Concentrations were calculated by comparing the peak area of sample peaks with calibration curves for peak areas of each standard compound. All concentrations are expressed as nanomoles per milligram of protein.

**Kinase assays.** AMPK from extracts was immunoprecipitated with a mixture of α1 and α2 antibodies, and AMPK activity in the immunoprecipitates was determined using the AMARA peptide assay as described previously (Hardie et al., 2000; Gadalla et al., 2004), except that, in the present study, the phosphorylation state of the native full-length protein was determined.

**Electrophysiological recordings.** Except for one series of experiments, slices (comprising hippocampus and overlying neocortex) were incubated for 3–8 h in standard aCSF or for 2 h in aCSF supplemented with 1 mM ribose/50 μM adenine (Rib/Ade) and 1–6 h in standard aCSF before being transferred to a recording chamber and fully submerged in aCSF at 33.4 ± 0.2°C. A flow rate of 6–7 ml/min was used. In the other series of experiments, slices were immediately transferred to the recording chamber during slice cutting to monitor the recovery of synaptic transmission in control and Rib/Ade-containing aCSF. A twisted bipolar Teflon-coated tungsten wire was placed to stimulate the Schaffer collateral/commisural pathway every 15 s, and field EPSPs (fEPSPs) were recorded from stratum radiatum in area CA1 of the hippocampus with a glass micro-electrode filled with aCSF (1 MΩ). The stimulus intensity was adjusted to 50–60% of that required to evoke a population spike. LTP was induced with tetanic stimulation (one train of 100 stimuli at 100 Hz) or with theta-burst (TBS) stimulation (0.5, 1, 2, or 3 × 10 trains of four stimuli at 100 Hz repeated at 200 ms intervals).

**Adenosine biosensors.** Adenosine and null microelectrode biosensors (50 μm diameter and 500 μm length) were purchased from Sarissa Biomedical Ltd. and were used to measure the real-time release of adenosine during LTP induction. The use of the sensors in hippocampal slices has been described previously (Frenguelli et al., 2003, 2007). The adenosine sensor relies on an enzyme cascade immobilized within a matrix on the surface of a platinum/iridium electrode to metabolize adenosine, thereby liberating H2O2, which is oxidized on the platinum/iridium electrode. This gives rise to an oxidation current proportional to the concentration of adenosine. The null sensor lacks enzymes and is used to establish the presence of any electroactive interferents. Both sensors were inserted into the stratum radiatum of the CA1 region of hippocampal slices between recording and stimulating electrodes. After insertion, slices were allowed to recover for 30–45 min before electrical stimulation for the recording of fEPSPs was started. After a stable fEPSP baseline of 15–20 min was collected, adenosine release was evoked with three TBS given 10 s apart. fEPSPs, adenosine and null sensor traces were recorded simultaneously. Thirty minutes after LTP induction, sensors were either taken out of the tissue or drugs were applied for 15–30 min before TBS was repeated. After each experiment, sensors were calibrated with 10 μM adenosine in the recording chamber. Because no nonspecific electroactive release could be detected on the null sensor, adenosine release was calculated without subtraction of the null trace, and the values are given as μM to reflect that the adenosine sensor measures adenosine and its metabolites (Frenguelli et al., 2007). To integrate the area under the curve of adenosine sensor traces, the baseline had to be set to 0, which was achieved by subtracting from the sensor trace a linear regression based on 5 min of baseline.

**Statistical analysis.** All values are expressed as mean ± SEM. For the electrophysiological and adenosine sensor measurements, n values refer...
to the number of slices per experimental condition, which for most cases is also equal to the number of animals used. Slices were used in duplicate for nucleotide extraction and in triplicate for protein extraction. In these cases, \( n \) values represent the number of animals used. For statistical analysis of more than two groups, one-way ANOVA with Bonferroni’s multiple comparison test was applied, whereas for comparisons between two independent groups, unpaired \( t \) tests were used. For comparison of the adenosine release before and after application of different drugs, a paired \( t \) test was applied. Calculations were performed with Prism 4; \( p \) values <0.05 were considered as statistically significant.

**Chemicals.** All HPLC standards, 1,1,2-trichloro-1,1,2-trifluoroethane (HPLC grade), EGTA, EDTA, sodium fluoride, sodium orthovanadate, sodium pyrophosphate, sodium azide, BSA, TBAHS, D-ribose, adenine, 8-cyclopentylethylphenoxyline (8-CPT), nitrobenzylthionosine (NBTH), di-pyridamole (DIPY), \( N^c \)-cyclopentyladenosine (\( N^c \)-CPA), sodium polyoxotungstate (POM-1), and the Bradford reagent were obtained from Sigma-Aldrich. Protease inhibitor cocktail tablets and ATP were from Roche. HPLC-grade methanol, perchloric acid, orthophosphoric acid, oxotungstate (POM-1), and the Bradford reagent were obtained from Thermo Fisher Scientific. Protein G Sepharose was from GE Healthcare. Pyridoxal phosphate-6-azophenyl-2',4'-disulfonic acid (PPADS) and forskolin were purchased from Ascent. Sheep antibodies against the \( a1 \) and \( a2 \) subunits of AMPK were described previously (Woods et al., 1996), and the antibody against the phosphorylated Thr-172 was from Cell Signaling Technologies.

**Results**

**Metabolic recovery after slice cutting**

**Recovery of adenine nucleotides**

To study the recovery of adenine nucleotides after slice preparation, HPLC analysis of slice extracts was performed on fresh slices immediately after cutting and after various incubation time points in aCSF (10 min to 5 h) at room temperature (22°C) and 34°C (Fig. 1A).

Immediately after cutting, ATP, ADP, and AMP were present in nearly equal amounts (6.0 ± 0.3, 4.0 ± 0.4, and 5.2 ± 0.8 nmol/mg protein, respectively; \( n = 7 \)) (Fig. 1B–D) (supplemental Table 1, available at www.jneurosci.org as supplemental material). ATP levels significantly increased after only 10 min incubation (10.7 ± 1.0 and 11.0 ± 0.9 nmol/mg protein at 22°C and 34°C, respectively; \( n = 7; p < 0.001 \), one-way ANOVA) with a concomitant decrease of ADP (1.9 ± 0.3 and 1.5 ± 0.1 nmol/mg protein at 22°C and 34°C, respectively; \( n = 7; p < 0.001 \), one-way ANOVA) and AMP (1.1 ± 0.3 and 0.5 ± 0.1 nmol/mg protein at 22°C and 34°C respectively; \( n = 7; p < 0.001 \), one-way ANOVA) levels. ATP degradation metabolites (IMP, adenosine, inosine, hypoxanthine, and xanthine) were all elevated at the time of cutting and together accounted for ~5 nmol/mg protein (supplemental Table 2, available at www.jneurosci.org as supplemental material). The levels of these metabolites declined after slice cutting and stabilized after 10–60 min of incubation.

After the initial recovery, ATP, ADP, and AMP levels did not significantly change during the incubation time points tested (up to 5 h incubation), and there were no significant differences between adenine nucleotides of slices kept at 22°C and 34°C (Fig. 1C and D). As a consequence of these complementary changes in individual nucleotides, the total adenine nucleotide pool (TAN = [ATP] + [ADP] + [AMP]) (Fig. 1E) did not significantly change when slices were transferred from the ice-cold cutting solution (15.2 ± 1.1 nmol/mg protein; \( n = 7 \)) into aCSF at 22°C (13.7 ± 1.4 nmol/mg protein; \( n = 7 \)) or 34°C (13.0 ± 1.1 nmol/mg protein; \( n = 7 \)), suggesting that most of the accumulated AMP is phosphorylated to ATP rather than dephosphorylated to adenosine (via cytosolic 5’-nucleotidase, EC 3.1.3.5), or deaminated to IMP (via AMP deaminase, EC 3.5.4.6.). The TAN pool remained stable over an incubation period of 5 h. Average TAN concentrations from all time points tested (10 min to 5 h) were 14.1 ± 0.3 nmol/mg protein in slices at 22°C and 15.4 ± 0.9 in slices at 34°C, with ATP accounting for ~85 and 89%, respectively.

**Recovery of energetic parameters and AMPK activity**

Two widely used measures of cellular energetic state are the adenylate energy charge, \( EC = (\text{[ATP]} + 0.5 \times \text{[ADP]})/\text{[TAN]} \) (Atkinson, 1968), which has a maximum value of 1 when all the adenine nucleotides are in the form of ATP and the ATP/AMP ratio.

Because of the nearly equal amounts of ATP, ADP, and AMP at the time of cutting, the \( EC \) was very low (0.54 ± 0.03; \( n = 7 \)) (Fig. 2A) (supplemental Table 3, available at www.jneurosci.org as supplemental material) but recovered significantly after only 10 min incubation at 22°C: \( 0.86 ± 0.019; n = 7; p < 0.001 \), one-way ANOVA) and 34°C (0.90 ± 0.007; \( n = 7; p < 0.001 \), one-way ANOVA). After 3 h the \( EC \) stabilized at 0.93 ± 0.003 for slices kept at 22°C and at 0.95 ± 0.002 for slices kept at 34°C, and, as for adenine nucleotides, there were no significant differences between slices at 22°C and 34°C.

The ATP/AMP ratio significantly recovered from time of cutting (1.4 ± 0.4; \( n = 7 \)) (Fig. 2B) (supplemental Table 3, available at www.jneurosci.org as supplemental material) after only 10 min in slices at 34°C (22.9 ± 2.7; \( n = 7; p < 0.001 \), one-way ANOVA), whereas it took ~30 min to recover in slices incubated at 22°C (20.1 ± 1.8; \( n = 6; p < 0.001 \), one-way ANOVA). Similar to the \( EC \), the ATP/AMP ratio stabilized after 3 h but with considerable differences between slices incubated at 22°C and at 34°C. At 22°C, the ATP/AMP ratio ranged between 35.2 and 38.0, whereas at 34°C, the ATP/AMP ratio was much higher between 63.5 and 64.2. The differences in the ATP/AMP ratio values between the two incubation temperatures became statistically significant after 30 min (\( n = 6–8; p < 0.01 \), one-way ANOVA) and remained statistically significant for the rest of the incubation period (\( n = 5–8; p < 0.001 \), one-way ANOVA).

The cellular ATP/AMP ratio is monitored by AMPK (EC 2.7.11.31), a key sensor and regulator of cellular energy metabolism (Hardie and Hawley, 2001; Hardie, 2007). AMPK is activated by phosphorylation of Thr172 by the upstream kinases LKB1 (Hawley et al., 2003) and calcium/calmodulin-dependent protein kinase kinase \( \beta \) (Hawley et al., 2005; Woods et al., 2005). In addition, an increase in cellular AMP provides both allosteric activation of the enzyme and protection of Thr172 from dephosphorylation, whereas both of these effects are antagonized by high intracellular ATP levels (Hardie et al., 2006; Sanders et al., 2007). Therefore, we investigated whether the lower ATP/AMP ratio in slices at 22°C was reflected by a higher AMPK activity.

AMPK activity (Fig. 2C) decreased from 0.066 ± 0.003 U/mg protein at time of cutting to 0.04 ± 0.001 U/mg protein after 30 min at 22°C (1.6-fold decrease; \( n = 3; p < 0.001 \), one-way ANOVA), by which time at 34°C AMPK activity had fallen 3.4-fold (\( n = 2 \)). After 3 h, AMPK was three times more active in slices at 22°C (0.03 ± 0.003 U/mg protein; \( n = 3 \)) than in slices at 34°C (0.01 ± 0.001 U/mg protein; \( n = 3; p < 0.001 \), one-way ANOVA). Western blots (Fig. 2D) showed that the ratio of phospho-AMPK/total AMPK decreased from 2.8 at time of cutting to 1.9 in slices at 22°C and to 1.1 in slices at 34°C after 3 h incubation (\( n = 2 \)) (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Likewise, the phospho/total ratio of a substrate, acetyl-CoA carboxylase (ACC), decreased from 1.6 at time of cutting to 1.0 in slices at 22°C and to 0.5 in slices at 34°C after 3 h incubation (Fig. 2D) (supplemental Fig. 1, available at
www.jneurosci.org as supplemental material). These observations suggest that, despite similarities between TAN pools and EC values between slices incubated at room temperature and more physiological temperatures, the ATP/AMP ratio can influence the activity of key intracellular enzymes with potentially important consequences for neuronal and glial properties.

**Basis of reduced TAN concentration in slices**

EC values of brain slices reported here (Fig. 2A) (supplemental Table 3, available at www.jneurosci.org as supplemental material) are comparable with those reported in vivo (supplemental references, available at www.jneurosci.org as supplemental material). However, absolute TAN levels here (Fig. 1E) (supplemental Table 1, available at www.jneurosci.org as supplemental material) and in the in vitro literature (supplemental references, available at www.jneurosci.org as supplemental material) are ~40–60% lower than published in vivo values for rat brain, which are typically ~33.6 ± 4.7 nmol/mg protein (arithmetic mean ± SD of all in vivo published data in supplemental references, available at www.jneurosci.org as supplemental material) (Fig. 3A). In our study, the loss of adenine nucleotides occurred either before or during slice preparation, because TAN levels were already 55% (~18.4 nmol/mg protein) lower at the time of cutting than published in vivo studies. We investigated several possible explanations for this observation.

The ischemic period leads to loss of diffusible ATP degradation products

The sum of ATP degradation metabolites (adenosine, inosine, hypoxanthine, xanthine, and IMP) at time of cutting was approximately ~5 nmol/mg protein (supplemental Table 2, available at www.jneurosci.org as supplemental material). Although, like ADP and AMP levels, these metabolites declined during the first 10–30 min of incubation, there was no corresponding rise in the TAN pool. Thus, they are likely to be lost from the tissue and thereby contribute to the reduced adenine nucleotide content of brain slices.

The tissue suffers from physical damage causing additional loss of adenine nucleotides

To establish whether the dissection of tissue associated with slicing caused additional loss of adenine nucleotides, we determined the TAN content immediately after decapitation/dissection in entire hippocampus and cortex (because slices in this study were composed of hippocampus and overlying neocortex). The TAN content of intact hippocampal and cortical tissue was higher than that in combined hippocampal/neocortical slices at 23.0 ± 2.1 and 28.9 ± 2.9 nmol/mg protein, respectively (Fig. 3A) (n = 5; p > 0.05 between hippocampal and cortical tissue, unpaired t test). The value for cortex is close to that reported in vivo (33.6 ± 4.7 nmol/mg protein) (Fig. 3A), but TAN levels in the hippocampus are lower than those reported in vivo. This suggests that the ischemic period during decapitation results in a loss of adenine nucleotides, especially in hippocampal tissue (~10 nmol/mg protein, ~29%), which additionally requires more physical and
potentially traumatic dissection for removal. In contrast, neocortex may undergo more rapid cooling when the brain is dropped into ice-cold aCSF, which may better preserve adenine nucleotides.

Figure 2. Differential influence of temperature on the recovery of energetic parameters and AMPK activity after slice cutting. Recovery of the tissue energy charge \((ATP + 0.5 ADP)/TAN\) is not influenced by temperature (A), whereas the ATP/AMP ratio is significantly higher at elevated temperature \((n = 3–8)\) (B). Accordingly, AMPK activity in brain slices, as measured by pseudo-substrate phosphorylation, is lower at higher incubation temperature, reflecting the higher ATP/AMP ratio \((n = 3\) except for 0.5 h, 34°C, \(n = 2\)). White squares, Slices at time of cutting (time 0); white triangles, slices incubated in aCSF at room temperature (22°C); black circles, slices incubated in aCSF at 34°C. C, Accordingly, AMPK activity in brain slices, as measured by pseudo-substrate phosphorylation, is lower at higher incubation temperature, reflecting the higher ATP/AMP ratio \((n = 3–8)\) except for 0.5 h, 34°C, \(n = 2\). White squares, Slices at time of cutting (time 0); white triangles, slices incubated in aCSF at room temperature (22°C); black circles, slices incubated in aCSF at 34°C. D, Confirmation of increased AMPK activity through Western blot analysis of increased phosphorylation of AMPK (p-AMPK) and a downstream target, ACC (p-ACC). Also shown are total AMPK and ACC at different durations and temperature of incubation in two separate sets of slices. All values are presented as mean ± SEM; \(n = 3–8\). C, Theoretical curves \((1 - \alpha)\) where \(\alpha = d/l\); dotted lines) to estimate the relative contribution of dead cut edges \((d)\) to the total tissue thickness of slices \((l)\), ranging from 100 to 2000 μm in 20 μm steps). Confirmation of increased AMPK activity through Western blot analysis of increased phosphorylation of AMPK (p-AMPK) and a downstream target, ACC (p-ACC). Also shown are total AMPK and ACC at different durations and temperature of incubation in two separate sets of slices. All values are presented as mean ± SEM. **p < 0.01, ***p < 0.001 compared between slices at 22°C and 34°C, one-way ANOVA with Bonferroni’s multiple comparison test. When no error bars can be seen, they are smaller than the symbol.

Figure 3. Tissue thickness and handling influence calculation of adenine nucleotide content of brain tissue. A, TAN content of whole hippocampus \((n = 5)\) and cortex \((n = 5)\) immediately after dissection and hippocampal \((n = 3)\) and neocortical slices \((n = 3)\) immediately after cutting. Note that whole tissue TAN levels are lower than reported in vivo values (dotted black line represents the arithmetic mean ± SD as shown by the gray area for all reported values from supplemental references, available at www.jneurosci.org as supplemental material), possibly reflecting increased handling/trauma, with only whole cortex approaching in vivo values. B, TAN levels of neocortical/hippocampal slices of varying thickness (200, 400, 800, 1200, 1500, and 3600 μm) immediately after cutting. All values in A and B are presented as mean ± SEM; \(n = 3–8\). C, Theoretical curves \((1 - \alpha)\) where \(\alpha = d/l\); dotted lines) to estimate the relative contribution of dead cut edges \((d)\) to the total tissue thickness of slices \((l)\), ranging from 100 to 2000 μm in 20 μm steps), assuming a total thickness for the two dead cut edges of 40 μm (light gray dotted line, 20 μm on each side of the slice), 70 μm (black dotted line, 35 μm on each side of the slice), or 100 μm (dark gray dotted line, 50 μm on each side of the slice). By normalizing the TAN levels obtained from slices at different thicknesses from B to the TAN values obtained for 3600 μm slices (plotted as black dots), we found that these normalized values fit the curve for a total dead cut edge layer of 70 μm (black dotted line), D, Theoretical curve for the ratio of the thickness of cut dead edges \((d)\) to the tissue thickness \((l)\) ranging from 100 to 3600 μm in 20 μm steps), showing that, with increasing slice thickness, the relative contribution of the thickness of the cut edges decreases, approaching a value of 0.99 (black circles).
The difference in TAN levels between cortex and hippocampus can also be seen in slices (Fig. 3A). Hippocampal slices had significantly lower TAN levels than cortical slices (16.4 ± 1.1 and 20.7 ± 0.7 nmol/mg protein, respectively; n = 3; p < 0.05, unpaired t test). These values were ~28% lower than the respective whole tissue values reported above and ~38% (cortex) to 50% (hippocampus) lower than reported in vivo values (Fig. 3A).

The dead layer on slice surfaces distorts adenine nucleotide measurements

**Empirical observations.** To test whether the protein content of dead slice edges (typically 35–50 μm) (Feig and Lipton, 1990; Siklós et al., 1997; Frenguelli et al., 2003) results in an underestimation of ATP in the viable core of the slice, we prepared neocortical/hippocampal slices of different thickness, thereby changing the ratio of dead to viable tissue. Because there were no significant changes in the TAN levels between slice cutting and 5 h incubation (Fig. 1E) (supplemental Table 1, available at www.neurosci.org as supplemental material) and to bypass the problem of a possible emerging nutrient-deprived core in very thick slices, the analysis was performed immediately after cutting.

The TAN content of slices, relative to the amount of protein, increased ~26% with increasing thickness from 15.2 ± 1.7 nmol/mg protein in a 200 μm slice to 20.6 ± 1.0 nmol/mg protein in a 1200 μm slice (Fig. 3B) (n = 5–8; p < 0.05, unpaired t test). There was no additional increase in the TAN levels in 1500 μm slices (20.3 ± 2.3 nmol/mg protein; n = 4), and we obtained a value of 21.1 ± 2.6 nmol/mg protein for 3600 μm slices (n = 3).

**Theoretical predictions.** To better understand the dependence of TAN content on slice thickness, we made the assumption that the TAN is proportional to the volume of the tissue (I^2) as defined by a unit of length l. If in a slice there is a layer of dead tissue of thickness d devoid of adenine nucleotides at either face of the slice, then the volume of tissue contributing to the TAN is I^2(l−d). We further assumed that d is constant and does not depend on the slice thickness and expressed d as a proportion of l (d = αl). The volume of tissue contributing to the TAN is thus I^2(l−lα). If we consider a unit of volume (i.e., l = 1), then a plot of 1−1−α against the normalized TAN for different slice thicknesses, assuming constant d, should fit our observed data and provide a theoretical estimate of the dead layer of tissue at either face of the slice. In Figure 3C, we have plotted theoretical curves for the relative contribution of a dead tissue layer (d) of 20 μm (40 μm in total), 35 μm (70 μm in total), and 50 μm (100 μm in total) on both slice edges to the total tissue thickness (l, from 100 to 2000 μm slices in 20 μm steps). By normalizing the measured TAN values in Figure 3B to the TAN value obtained for 3600 μm slices (21.1 nmol/mg protein) and plotting it on the same graph, we observed a very good fit of our measured values to the theoretical curve obtained for an estimated total dead cut edge layers of 70 μm, or 35 μm for each edge (Fig. 3CD, black dotted lines). We previously reported a value of 35 μm as an estimate of the dead slice layer based on histological assessment of 400 μm slices (Frenguelli et al., 2003), revealing a remarkable degree of consistency between our experimental observation and our theoretical model.

With this curve (ranging from 100 to 3600 μm in 20 μm steps) (Fig. 3D), an asymptotic value is approached at a tissue thickness of 3600 μm (1−α = 0.980), suggesting that the dead cut edges account for only 2% of the whole tissue thickness. Therefore, assuming a maximal TAN value of 21.1 nmol/mg protein (3600 μm slices) in slices, we might underestimate the TAN content of the viable core tissue in a 400 μm slice (17.1 ± 1.5 nmol/mg protein) by ~4 nmol/mg protein, ~19%. Nonetheless, when corrected for this amount, slice TAN levels in 400 μm neocortical/hippocampal slices remain ~37% (~12.5 nmol/mg) lower than reported in vivo values (33.6 ± 4.7 nmol/mg protein as shown by the gray area for the mean ± SD in Fig. 3B). Hence, this difference is likely attributable to the loss of adenine nucleotides and precursors during the ischemia and physical trauma associated with slice preparation.

**Supplementation of aCSF with adenine nucleotide precursors improves cellular ATP levels**

In vivo cerebral TAN or ATP levels recover after brief periods of ischemia (~1–5 min) to pre-ischemic values after ~60–90 min reperfusion (Ljunggren et al., 1974; Kobayashi et al., 1977; Nowak et al., 1985). However, there was no significant increase in TAN levels in slices over a 5 h incubation period (Fig. 1E). This might be attributable to a lack of purine precursor metabolites in the aCSF, which might otherwise be used to restore tissue ATP levels via purine salvage or de novo synthesis.

Because two key components of the purine salvage pathway, which is believed to predominate in brain (Gerlach et al., 1971; Mascia et al., 2000; Barsotti and Ipata, 2002), are adenine and D-ribose (Fig. 4A), we tested these compounds in brain slices. Incubating slices in 50 mM Ade and 1 mM Rib resulted in tissue levels of Ade reaching a maximum after 1 h incubation (0.92 ± 0.07 nmol/mg protein), with no additional increase after 3 h (0.95 ± 0.04 nmol/mg protein; n = 3–5) (data not shown). Interestingly, the uptake of Ade was facilitated by 1 mM Rib (0.56 ± 0.09 nmol/mg protein after 3 h incubation in 50 mM Ade alone compared with 0.95 ± 0.04 nmol/mg protein after 3 h incubation in 1 mM Rib/50 μM Ade; p < 0.05, one-way ANOVA), with no additional increase observed with higher Rib concentrations (0.83 ± 0.06 nmol/mg protein after 3 h incubation with 10 mM Rib/50 μM Ade, n = 3) (data not shown).

To test whether Ade and Rib could be used by the purine salvage pathway to restore adenine nucleotide levels in brain slices, we incubated freshly cut slices in aCSF supplemented with 1 mM Rib and 50 μM Ade (Fig. 4). The TAN content in slices incubated with Rib/Ade increased to 25.8 ± 0.7 nmol/mg protein after 3 h incubation (compared with 19.1 ± 1.2 nmol/mg protein in slices incubated in standard aCSF; n = 3–5; p < 0.01, one-way ANOVA) (Fig. 4B), with ATP accounting for ~92%. When corrected for the influence of the protein content of the dead slice edges (~4 nmol/mg protein) (Fig. 4B), these values (~30 nmol/mg protein) are close to the values reported for in vivo tissue (33.6 ± 4.7 nmol/mg) (Fig. 4B). The elevation of tissue TAN and ATP levels by Rib/Ade did not significantly impact on the EC, which stabilized at 0.96 ± 0.001, and the ATP/AMP ratio, which reached an asymptotic value after 3 h at 141.7 ± 17.5 compared with 94.9 ± 23.5 for slices incubated in standard aCSF (n = 3–5; p > 0.05, one-way ANOVA) (Fig. 4C).

Lower Rib concentrations (500 μmol) were not as effective in increasing TAN levels and higher Rib concentrations (2.5–10 mM) did not further increase the TAN content of slices (data not shown). Furthermore, Ade (50 μmol) on its own, as well as higher Rib concentrations (2.5 or 10 mM) on its own did not significantly increase the TAN content in slices (supplemental Table 4, available at www.neurosci.org as supplemental material). This suggests that both metabolites are needed for effective conversion to adenine nucleotides during a 3 h incubation period.

To establish whether these elevated levels of TAN and ATP persisted when Rib/Ade was removed, we incubated slices in standard aCSF for 2 h after 3 h in Rib/Ade. During this time, tissue adenine content decreased back to baseline (0.1 ± 0.05
These data suggest that the full recovery of slice ATP levels is limited by the lack of ATP precursors in the aCSF. In addition, these data also indicate that providing ATP precursors in the form of Rib/Ade allows the viable core of brain slices to restore ATP levels to values close to those reported in vivo.

Electrophysiological properties of slices incubated in Rib/Ade

To establish whether the higher ATP and TAN levels in slices incubated in Rib/Ade would alter the electrophysiological properties of brain slices, we performed extracellular recordings from the CA1 region of hippocampal slices. Input–output curves, paired-pulse facilitation, and LTP were compared between slices incubated for 3–8 h in standard aCSF and slices incubated for 2 h in 1 mM Rib and 50 μM Ade then for 1–6 h in standard aCSF to wash these agents out of the tissue. A 2 h incubation period in Rib/Ade was chosen, because slice TAN levels reached an asymptotic value at that time (25.4 ± 2.3 nmol/mg protein), with no additional increase after 3 h (25.8 ± 0.6 nmol/mg protein; n = 3–5) (data not shown).

Basal synaptic transmission is normal in Rib/Ade-treated slices

The recovery of synaptic transmission after slice cutting was not different between standard and Rib/Ade-treated slices (supplemental Fig. 2, available at www.jneurosci.org as supplemental material). Furthermore, in a separate series of slices, there was no significant difference in input–output curves (Fig. 5A; n = 15–16) and paired-pulse ratios (Fig. 5B; n = 18–22) between the two set of slices (p > 0.05, one-way ANOVA). Likewise, 50 μM Ade on its own (Fig. 5C; n = 4) or in combination with 1 mM Rib (Fig. 5D; n = 6) did not change paired-pulse ratios when acutely applied to slices. This suggests that, under conditions of low-frequency stimulation of afferent fibers, the enhanced tissue ATP levels in Rib/Ade-treated slices is not being released to form adenosine in the extracellular space, which would, via inhibitory A1Rs, inhibit glutamate release and raise the paired-pulse facilitation ratio. These negative results suggest that the activation A1Rs and basal handling of adenosine is normal between standard and Rib/Ade-treated slices. To test this directly, we applied the selective A1R agonist N6-CPA (10 μM) (Gadalla et al., 2004) or the adenosine uptake inhibitors NBPTI (5 μM)/DIPY (10 μM) (Frequelli et al., 2007; Etherington et al., 2009) to both sets of slices (Fig. 5E,F). The concentrations chosen were submaximal for complete depression of the iEPSP to avoid a “floor effect” obscuring potential differences between the two sets of slices. Furthermore, we have shown previously that NBPTI/DIPY causes a depression of the iEPSP that can be reversed with A1R antagonists (Pearson et al., 2001) and have demonstrated the increase in extracellular adenosine directly with adenosine biosensors (Frequelli et al., 2007; Etherington et al., 2009).

The rate and extent of iEPSP depression after a 15 min application of N6-CPA was the same in control (50.0 ± 1.6%; n = 5) and Rib/Ade-treated (50.4 ± 1.6%; n = 5; p > 0.05, unpaired t test) slices (Fig. 5E). Likewise, the application NBPTI/DIPY for 40 min resulted in the same rate and amount of depression in both sets of slices (40 ± 5.9% for standard slices; 43 ± 3.3% for Rib/Ade treated slices; n = 4; p > 0.05, unpaired t test) (Fig. 5F).

These data suggest that Rib/Ade pretreatment does not influence the sensitivity of the A1R to agonists, nor is the activity of equilibrative adenosine transporters affected.

Furthermore, the fact that acute application of Ade did not change paired-pulse ratios (Fig. 5C,D), shows that the recently described Gαi-protein-coupled adenine receptor (Bender et al., 2002; von Kügelgen et al., 2008), if present in the hippocampus,
does not have any presynaptic effects on neurotransmitter release. However, to further exclude the possibility of differences in cAMP formation between Rib/Ade-treated slices and slices incubated in standard aCSF, we applied 50 μM forskolin to both sets of slices and compared the increase in fEPSP slopes (Fig. 5G,H). There was no significant difference in forskolin-induced potentiation between Rib/Ade-treated slices and slices incubated in standard aCSF (Fig. 5G) (147.4 ± 9.8% in standard slices and 159.1 ± 12.9% in Rib/Ade-treated slices; n = 3–4; p < 0.05 compared with baseline before application of forskolin, p > 0.05 between standard slices and Rib/Ade-treated slices at 20 min after application of forskolin, one-way ANOVA). Paired-pulse facilitation (50 ms interpulse interval) was similarly affected by forskolin in standard slices and Rib/Ade-treated slices (Fig. 5H) (n = 3–4; p > 0.05 between standard slices and Rib/Ade-treated slices, unpaired t test). This suggests that adenylyl cyclase activation and cAMP production is not impaired in Rib/Ade-treated slices.

**Long-term potentiation is impaired in Rib/Ade-treated slices**

Slices incubated in standard aCSF showed robust LTP 55–60 min after tetanic stimulation (one train of 100 shocks at 100 Hz; 135 ± 5.8% of baseline; n = 9; p < 0.001 compared with 5 min baseline before tetanic stimulation, one-way ANOVA) (Fig. 6A). However, LTP in slices incubated for 2 h in 1 mM Rib and 50 μM Ade decayed back to baseline 60 min after tetanic stimulation (108 ± 4.9% of baseline; n = 11; p > 0.05 compared with 5 min baseline before tetanic stimulation, one-way ANOVA) and was significantly lower in amplitude than LTP in control slices incubated in standard aCSF (p < 0.001 from 55 to 60 min after LTP induction, one-way ANOVA). Stable recordings could be achieved in Rib/Ade-treated slices over the same time period (94 ± 6.6% of baseline at 75 min; n = 3) (data not shown), which argues against baseline drift as being the cause for the observed decay in LTP. In contrast, acute application of Ade or Rib alone, or in combination, did not impair tetanus-induced LTP (133 ± 5% of baseline for standard slices, 131 ± 13% after acute application of 50 μM Ade, 140 ± 9% after acute application of 1 mM Rib, and 131 ± 8% after acute application of 1 mM Rib/50 μM Ade at 60 min after LTP induction; n = 4–5) (data not shown), implying a requirement for uptake and intracellular conversion to adenine nucleotides.

**Figure 5.** Basal synaptic transmission, adenosine A1 receptor activation, and adenosine uptake are not different between slices incubated in standard aCSF and slices treated for 2 h in 1 mM Rib and 50 μM Ade. Input–output curves (n = 15–16) (A) and paired-pulse ratios (n = 18–22) (B) for slices incubated in standard aCSF (black circles) and slices treated for 2 h with Rib/Ade-supplemented aCSF (gray circles). Insets are representative fEPSPs from 10–300 μA and at 50 ms interpulse interval (B) for controls and Rib/Ade-treated slices. C,D, Paired-pulse ratios for slices incubated in standard aCSF before (black circles) and after acute application of 50 μM Ade (C, white circles, n = 4) or 50 μM Ade and 1 mM Rib (D, white circles, n = 6). Insets are representative fEPSPs at 50 ms interpulse interval before (black traces) and after application of Ade or Rib/Ade (dotted gray traces). There was no difference in the rate or magnitude of fEPSP depression in response to the selective A1 receptor antagonist N6-CPA (10 μM, n = 5) (E) or the combination of the adenosine uptake inhibitors NBTI (5 μM)/DIPY (10 μM, n = 4) in standard slices or Rib/Ade-treated slices (F). G, Forskolin-induced potentiation in standard slices (black circles) and Rib/Ade-treated slices (gray circles). Forskolin at 50 μM was applied to slices for 20 min and no differences were observed in the amount of potentiation (147 ± 5.8% in standard slices and 159.1 ± 12.9% in Rib/Ade treated slices; n = 3–4) or the decrease in paired-pulse facilitation (p = 0.2) at a flow rate of 7–8 ml/min after a 2 h recovery period from slice cutting. All values are presented as mean ± SEM. No significant differences observed between standard and Rib/Ade-treated slices with unpaired t tests or one-way ANOVA with Bonferroni’s multiple comparison test. When no error bars can be seen, they are smaller than the symbol.
A potential reason for the impaired LTP stabilization in Rib/Ade-treated slices might be that, because of the higher TAN pool, enhanced synaptic activity during tetanic stimulation causes the activity-dependent release of ATP and/or its metabolite adenosine. To test this, we incubated slices in the ATP P2 receptor antagonist PPADS (10 μM) and A1R antagonist 8-CPT (1 μM) before the induction of LTP. Preincubation with PPADS did not prevent the decay of LTP in Rib/Ade-treated slices (108 ± 15% of baseline 60 min after LTP induction; n = 4) (data not shown). These observations suggest that, if ATP is being released during tetanic stimulation, it is not directly responsible via P2y receptors for the impairment of LTP.

To test for a role for adenosine A1 receptors, we applied 8-CPT (1 μM), a selective A1R antagonist, to slices that had either been pretreated with Rib/Ade or incubated in standard aCSF. Ten minutes after acute application of 8-CPT, fEPSPs had increased to 121.4 ± 0.7% in standard slices (n = 6) (data not shown) and to 124.1 ± 0.5% in Rib/Ade-treated slices (n = 8) (data not shown) and was associated with a similar decrease in PPF in both (data not shown). Both these changes in synaptic transmission are indicative of the removal of a basal adenosine A1R-dependent inhibitory tone. The fact that the changes were similar between the two conditions argues against the possibility of an increased basal adenosine tone in Rib/Ade-treated slices. This is consistent with the following observations: (1) similar tissue adenosine levels in both sets of slices after a 3 h incubation (standard slices, 0.05 ± 0.004 nmol/mg protein, n = 3; Rib/Ade-treated slices, 0.04 ± 0.002; n = 3; p > 0.05, unpaired t test) (data not shown), (2) normal basal transmission and paired-pulse facilitation in Rib/Ade-treated slices (Fig. 5A–D), (3) identical effects of A1R activation and uptake inhibition (Fig. 5E,F), and (4) equal effects of forskolin (Fig. 5G,H).

Having established that the basal handling and effects of adenosine were similar in control and Rib/Ade-treated slices, we next examined the effect of the A1R antagonist on tetanus-induced LTP with or without previous treatment with Rib/Ade. LTP was induced when a stable fEPSP baseline was collected, ~30–40 min after application of 8-CPT. Both sets of slices showed robust LTP in the presence of an increased basal adenosine tone in Rib/Ade-treated slices. (Fig. 6A–D) (143 ± 6.0% in standard slices and 154 ± 6.7% in Rib/Ade-treated slices; n = 5–6; p < 0.001 compared with 5 min baseline before tetanic stimulation, p > 0.05 between standard slices and Rib/Ade-treated slices 55–60 min after tetanic stimulation, one-way ANOVA). This suggests that the activity-dependent release of adenosine contributes to the impairment of LTP induction in Rib/Ade-treated slices.

We hypothesized that this deficit in LTP induction represented a raising of the threshold for LTP by the activity-dependent accumulation of extracellular adenosine and activation of inhibitory A1Rs. This hypothesis predicted that stronger activation of postsynaptic neurons should overcome
Control and Rib/Ade-treated slices were stimulated with two TBS at 5 min intervals (2 × 10 trains of four pulses at 100 Hz with 200 ms intervals, i.e., 80 pulses in total). This protocol resulted in robust LTP in both standard and Rib/Ade-treated slices 55–60 min after TBS (Fig. 6C) (150.2 ± 7.6% in standard slices and 140.0 ± 7.1% in Rib/Ade-treated slices; n = 6–7; p < 0.001 compared with 5 min baseline before TBS, p > 0.05 between standard slices and Rib/Ade-treated slices at 55–60 min after TBS, one-way ANOVA). One TBS (40 pulses) resulted in smaller LTP 55–60 min after TBS and showed little difference between standard and Rib/Ade-treated slices (Fig. 6D) (137.7 ± 9.5% in standard slices and 126.2 ± 5.0% in Rib/Ade-treated slices; n = 6; p < 0.01 for standard slices and p < 0.05 for Rib/Ade-treated slices compared with 5 min baseline before TBS, p > 0.05 between standard slices and Rib/Ade-treated slices at 55–60 min after TBS, one-way ANOVA).

To confirm that TBS resulted in greater depolarization compared with tetanic stimulation, we measured the area associated with each pulse induced by tetanic stimulation (100 pulses) and TBS (40 pulses) in a manner similar to that described recently (Chen et al., 2010). A comparison of the normalized cumulative area evoked by each pulse in a tetanus and TBS (supplemental Fig. 3A, available at www.jneurosci.org as supplemental material) revealed a dramatic difference between the two: in a tetanus, most of the depolarization had occurred within the first 20 pulses, whereas during TBS, the depolarization increased almost linearly during the 40 pulse train.

Furthermore, during a tetanus, there was evidence of an influence of Rib/Ade in causing fatigue of transmission during the later stages of the train (>20 pulses) (supplemental Fig. 3A, available at www.jneurosci.org as supplemental material). This was prevented in slices treated with 8-CPT (supplemental Fig. 3B, available at www.jneurosci.org as supplemental material) and is consistent with a gradual synaptic accumulation of adenosine. Indeed, the apparent enhancement in 8-CPT/Rib/Ade-treated slices may reflect an action of adenosine on facilitatory adenosine receptors (Chen et al., 2010).

To establish whether the released adenosine arose from direct release of adenosine or from extracellular degradation of ATP, we used adenosine uptake inhibitors (NBPT/DIPY) and POM-1, a noncompetitive inhibitor of ectonucleotidases (Muller et al., 2006; Wall et al., 2008). To assess the effect of these drugs on adenosine release, TBS-induced adenosine release was evoked twice (45–60 min apart): the first TBS was in control aCSF and slices treated with Rib/Ade (Fig. 7B). Rib/Ade-treated slices released significantly more adenosine during TBS as measured by integrating the area under the adenosine signal at the start of TBS to 5 min after stimulation (Fig. 7B) (0.64 ± 0.1 μM*min in slices incubated in standard aCSF and 1.98 ± 0.1 μM*min in Rib/Ade-treated slices; n = 3; p < 0.01, unpaired t test). Despite this greater release of adenosine, we could not observe any differences in LTP (Fig. 7A) between the two sets of slices as measured at 30 min after TBS. This is likely attributable to the fact that the strong stimulation protocol (Fig. 6C, D) overcame the inhibitory effects of A1R.

These results suggest that higher intracellular TAN or ATP levels result in increased activity-dependent adenosine release during periods of strong electrical stimulation, which can modulate the induction threshold for LTP.

Mechanism of activity-dependent adenosine release
To establish whether the released adenosine arose from direct release of adenosine or from extracellular degradation of ATP, we used adenosine uptake inhibitors (NBPT/DIPY) and POM-1, a noncompetitive inhibitor of ectonucleotidases (Muller et al., 2006; Wall et al., 2008). To assess the effect of these drugs on adenosine release, TBS-induced adenosine release was evoked twice (45–60 min apart): the first TBS was in control aCSF (in either standard slices or Rib/Ade-treated slices), and the second was in the presence of NBPT (5 μM)/DIPY (10 μM) or POM-1 (100 μM), which were applied 30 min after the initial TBS. Repeating TBS twice within 45–60 min did not affect adenosine...
release (supplemental Fig. 4A, available at www.jneurosci.org as supplemental material). If adenosine was released directly, we would expect the transport inhibitors to reduce TBS-induced adenosine accumulation as they represent a major efflux pathway for adenosine into the extracellular space (Baldwin et al., 2004). If extracellular adenosine arose from the metabolism of ATP, POM-1 should reduce TBS-induced adenosine release.

The effectiveness of POM-1 to inhibit ATP breakdown in hippocampal brain slices was assessed by inserting adenosine biosensors into slices and measuring adenosine production after exogenous ATP application in the presence and absence of POM-1 (Wall et al., 2008). POM-1 caused a time-dependent inhibition of ATP breakdown: in the absence of POM-1, 50 μM ATP yielded 3.5 ± 0.2 μM adenosine. After application of POM-1 for 5 min, ATP breakdown was inhibited by 30 ± 3% (2.5 ± 0.1 μM; n = 3; p > 0.05 compared with initial response, one-way ANOVA), whereas after 15 min, the metabolism of ATP was inhibited by 42 ± 2% (2 ± 0.2 μM; n = 3; p < 0.05 compared with initial response, one-way ANOVA) (supplemental Fig. 4B,C, available at www.jneurosci.org as supplemental material). We therefore decided to wait 15 min after POM-1 application to study the effect of POM-1 on TBS-induced adenosine release.

In the presence of POM-1, TBS-induced adenosine release was variable but showed no evidence of inhibition of adenosine release. In fact, the reverse was observed: POM-1 seemed to increase TBS-induced adenosine release in both standard and Rib/Ade slices (supplemental Fig. 4A, available at www.jneurosci.org as supplemental material). This could represent an off-target effect, as we have suggested exist in the use of POM-1 (Wall et al., 2008), or could reflect a previously described ATP-mediated facilitation of adenosine release via the activation of ATP P2 receptors (Almeida et al., 2003). To address this, we incubated Rib/Ade-treated slices with the P2 antagonist PPADS (10 μM) for 10 min before TBS. PPADS had no significant effect on adenosine release in Rib/Ade-treated slices (2.0 ± 0.41 μM*min for Rib/Ade-treated slices; n = 10; and 2.0 ± 0.61 μM*min for Rib/Ade-treated slices in the presence of PPADS; n = 3; p > 0.05, unpaired t test), and indeed did not affect the increased TBS-induced adenosine release in the presence of POM-1 (2.8 ± 0.45 μM*min for Rib/Ade-treated slices after POM-1 application; n = 4; and 2.5 and 3.3 μM*min in two Rib/Ade-treated slices in the presence of POM-1 and PPADS) (supplemental Fig. 4A, available at www.jneurosci.org as supplemental material). These negative results with the ectonucleotidase inhibitor POM-1 argue against an appreciable release of ATP and extracellular conversion to adenosine.

In contrast, application of NBTI/DIPY for 30 min resulted in a 50% reduction in TBS-induced adenosine release in both sets of slices (supplemental Fig. 4A, available at www.jneurosci.org as supplemental material) (50.3 ± 17.0% for control slices, n = 4; p = 0.06; 49.2 ± 11.0% for Rib/Ade-treated slices, n = 3, p = 0.04, paired t test), suggesting a role of equilibrative adenosine transporters in the release of adenosine in response to high-frequency stimulation of afferent fibers.

**Discussion**

Despite the importance and widespread use of brain slices as models of the mammalian CNS, criticisms remain regarding their metabolic integrity. Our aim was to address this issue to study (1) the energetic recovery of brain slices, (2) the reasons for reduced levels of ATP, (3) the possibility of improving cellular ATP, and (4) the functional consequences of raising tissue ATP levels.

**Energetic recovery after slice preparation**

In accordance with previous findings (Fredholm et al., 1984; Whittingham et al., 1984a,b), our results show that adenine nucleotide levels in brain slices recover quickly and remain stable for at least 5 h, independently of the incubation temperature. Likewise, the EC and ATP/AMP ratio show a rapid recovery after slice preparation, but it takes 3 h until they stabilize, well beyond the time conventionally allowed for slices to recover. Provided with an adequate supply of nutrients, it is likely that both interface and submerged slices will recover similarly in terms of adenine nucleotides and energetic parameters. Indeed, TAN and adenosine levels for interface hippocampal slices (~10.5 nmol/mg protein and 40 pmol/mg protein, respectively) (Fredholm et al., 1984) are not different from our results (~9.8 nmol/mg protein and 50 pmol/mg protein, respectively).

However, it is worth noting that other metabolites, such as cGMP and cAMP, lactate, or phosphocreatine, also require 1–3 h to achieve a steady state (Whittingham et al., 1984b). Likewise, the phosphorylation status of proteins involved in synaptic plasticity, such as GluA1, ERK2, and MEK1/2, changes during the first 3 h of incubation (Ho et al., 2004), whereas a recovery period of 4 h has been suggested for achieving stable long-term recordings of LTP in brain slices (Sajikumar and Frey, 2004; Sajikumar et al., 2005; Redondo et al., 2010).
Temperature dependence of the ATP/AMP ratio and AMPK activity

Our results indicate that slices maintained at 34°C have a significantly higher ATP/AMP ratio compared with slices at room temperature. This is likely attributable to the activity of adenylate kinase (EC 2.7.4.3; 2 ADP ↔ ATP + AMP), which is greater at temperatures above 32°C (Sheng et al., 1999; Lu and Wang, 2008). Accordingly, slices incubated at 22°C had a lower ATP/AMP ratio and showed a threefold higher AMPK activity than slices maintained at 34°C. This translated into increased phosphorylation of a downstream target, ACC, and suggests that other downstream targets are likely to be similarly affected.

Of the known AMPK targets relevant to synaptic physiology, AMPK phosphorylates GABA_A receptors (Hardie and Franguelli, 2007; Kuramoto et al., 2007) and calcium-activated potassium channels (Wyatt et al., 2007). Furthermore, a proteomic screen revealed 12 brain-specific downstream targets of AMPK, including synapsin I and PACSIN1, further suggesting a role for AMPK in regulating synaptic activity (Tuerk et al., 2007). Indeed, enhancing AMPK activity inhibits long-lasting LTP (Potter et al., 2010). Hence, incubating slices at elevated temperatures will more closely replicate metabolism in vivo with respect to the activity of enzymes and properties of synaptic transmission.

Basis of reduced tissue ATP content in brain slices

As seen in this and many other reports (McIlwain, 1952; Thomas, 1957; Kass and Lipton, 1982; Fredholm et al., 1984; Whittingham et al., 1989; Paschen and Djuricic, 1995; Milusheva et al., 1996), brain slices have ~40–60% lower ATP and TAN levels than the in vivo brain (Supplemental References). Our results suggest that this is attributable to the loss of adenine nucleotides and/or their metabolites, especially in hippocampal tissue, during slice preparation. However, incubation of slices in Rib/Ade-supplemented aCSF allowed the tissue to appreciably increase TAN levels. When corrected for the influence of the dead slice edges, which we could model accurately and which agreed with histological estimates we made previously (Franguelli et al., 2003), TAN levels were within the range reported in vivo. This suggests that the recovery of ATP levels in brain slices is limited by the lack of precursors in the aCSF and does not necessarily reflect an intrinsic metabolic handicap.

Implications of improving the tissue ATP content in brain slices

Although Rib/Ade restored tissue nucleotide levels close to those observed in vivo, this did not have a bearing on basal synaptic transmission, paired-pulse facilitation, or the tonic handling and effects of extracellular adenine. Instead, Rib/Ade inhibited LTP after tetanic stimulation and weak TBS. That the application of an A1R antagonist reversed the fatigue of the fEPSP during the tetanus and prevented the decline of tetanus-induced LTP in Rib/Ade-treated slices suggests that the higher TAN levels resulted in greater activity-dependent adenosine release, thereby preventing the stable expression of LTP. This suggests that endogenous adenosine exerts an inhibitory influence on LTP induction (de Mendonça and Ribeiro, 1994; Forghani and Krnjevic, 1995; Fujii et al., 2000; Rex et al., 2005), especially when used with weak stimulation protocols (Arai and Lynch, 1992; de Mendonça and Ribeiro, 2000). Interestingly, the threshold for adenosine-dependent regulation of TBS, based on the facilitatory actions of an A1R antagonist, was 20 pulses (Arai and Lynch, 1992; de Mendonça and Ribeiro, 2000), consistent with our predictions based on tetanic cumulative depolarizations and experiments with Rib/Ade-treated slices. These observations and our own results point toward an adenosine A1R-dependent regulation of LTP, which is influenced by the levels of intracellular adenine nucleotides.

However, our analysis of the TBS stimulus trains revealed no A1R-dependent fatigue of the fEPSP. This may reflect the fact that the 200 ms burst spacing may allow time for the removal of extracellular adenosine between stimulus trains through metabolism, reuptake, or diffusion. Thus, to the two known actions of TBS that make it an effective and naturalistic stimulus for LTP induction, maximizing both postsynaptic depolarization and GABAergic fatigue, we may now potentially add a third: minimizing the intraburst synaptic accumulation of extracellular adenosine.

Using adenosine biosensors, we were able to detect in real time the release of adenosine during TBS. Rib/Ade-treated slices released significantly more adenosine, consistent with the availability of a greater precursor pool of ATP. To establish whether ATP or adenosine was released in response to high-frequency stimulation, we used the noncompetitive ectonucleotide inhibitor POM-1 (Müller et al., 2006; Wall et al., 2008) and the equilibrative nucleoside transporter (ENT) inhibitors DIPY/NBTI (Franguelli et al., 2007; Etherington et al., 2009). POM-1 failed to reduce, and indeed facilitated, TBS-induced adenosine release. This is unlikely to be attributable to ATP P2 receptor-mediated facilitation of adenosine release reported by others (Almeida et al., 2003) because the facilitation was not affected by the P2 antagonist PPADS. Instead, this facilitation may involve nonspecific actions of POM-1 (Wall et al., 2008) or could potentially involve ATP heteroexchange with adenosine (Sperläch et al., 2003), which is insensitive to P2 receptor antagonists but sensitive to ENT inhibition. Accordingly, DIPY/NBTI caused a 50% reduction in TBS-induced adenosine release, which is consistent with the direct release of adenosine during high-frequency stimulation (Wall and Dale, 2008; Klyuch et al., 2011).

In a wider context, the reduced tissue ATP levels observed after cerebral ischemia in vivo may, via reduced extracellular adenosine and reduced activation of the anticonvulsant A1R (Boison and Stewart, 2009; Dale and Franguelli, 2009), contribute to the development of post-ischemic epilepsy (Camilo and Goldstein, 2004; Kadam et al., 2010). Indeed, the influence of intracellular ATP on extracellular adenosine and neuronal excitability has been described recently (Kawamura et al., 2010) and may be the basis for the reduced incidence of seizures during a ketogenic diet (Masino and Geiger, 2008). Accordingly, elevation of tissue ATP levels with Rib/Ade may be of value in the post-ischemic brain. In fact, Rib has been used to improve post-ischemic cardiac function in vitro, in vivo, and in humans (Zimmer, 1998; Omran et al., 2003; Shecterle et al., 2010). Although Ade has to be administered with a xanthine oxidase inhibitor (Watts et al., 1974; Simmonds, 1986) to prevent its conversion to an insoluble metabolite, this may be beneficial (Phillis et al., 1995) because it would prevent the formation of nonsalvageable xanthine, thereby providing greater substrates for the purine salvage pathway in the post-ischemic brain. Thus, Rib/Ade may be of value in restoring ATP levels and adenosine release after brain injury.

In summary, the data presented address long-standing issues in the use of brain slices as in vitro models for the mammalian CNS. We confirm the long-held view that tissue adenine nucleotides are ~50% of the values reported in vivo but demonstrate that this is an underestimate (by ~20%) because of the contribution of damaged slice edges. Moreover, we show that slices have
an appreciable capacity, through the purine salvage pathway, to restore and maintain tissue ATP levels close to in vivo levels when presented with the ATP precursors Ade and Rib. The physiological consequences of elevated tissue ATP levels are in the greater activity-dependent release of adenosine and the raising of the threshold for the induction of LTP.

References


