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Methodological standards for in vitro models of epilepsy and epileptic seizures. A TASK1-WG4 report of the AES/ILAE Translational Task Force of the ILAE

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Conflict disclosures

This report was written by experts selected by the International League Against Epilepsy (ILAE) and the American Epilepsy Society (AES) and was approved for publication by the ILAE and the AES. Opinions expressed by the authors, however, do not necessarily represent the policy or position of the ILAE or the AES. Reference to websites, products or systems that are being used for in vitro electrophysiological studies was based on the resources known to the co-authors of this manuscript and is done only for informational purposes. The AES/ILAE Translational Task Force of the ILAE is a non-profit society that does not preferentially endorse certain of these resources, but it is the readers' responsibility to determine the appropriateness of these resources for their specific intended experimental purposes.

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Ethical publication statement

We confirm that we have read the Journal's position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

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Summary

In vitro preparations are a powerful tool to explore the mechanisms and processes underlying epileptogenesis and ictogenesis. In this review, we critically review the numerous *in vitro* methodologies utilized in epilepsy research. We provide support for the inclusion of detailed descriptions of techniques, including often ignored parameters with unpredictable yet significant effects on study reproducibility and outcomes. In addition, we explore how recent developments in brain slice preparation relate to their use as models of epileptic activity.

Keywords

brain slice preparation; electrophysiological recording methods; recording solution composition; in vitro models of seizures; animal selection and killing

Introduction

In vitro preparations are a valuable and useful means of studying epilepsy and epileptic seizures¹. These preparations can be obtained from almost any species. The most widely used preparation in studies of epileptiform activity is the mammalian brain slice. These thin brain slices can either be used acutely (the acute brain slice preparation) or after preservation in culture in an incubator over a period of days or weeks (organotypic brain slice preparation). In addition, relatively intact brain preparations are of considerable value including whole hippocampus² or whole brain models^{3,4}. *In vitro* epileptiform activity can be evoked by performing ionic or pharmacological manipulations on preparations from naïve animals, by using tissue from animals who have experienced an epileptogenic *in vivo* insult, or by using genetic models of epilepsy with an *in vitro* phenotype. In this review, we do not compare or evaluate the multitude of *in vitro* models that exist. Rather we focus on the methodological details related to the establishment and utilization of *in vitro* preparations for studying seizures and epilepsy.

Animal selection and method of sacrifice

Choice of species, strain, age, and sex

Species—*In vitro* epileptiform activity has been elicited from brain preparations derived from multiple mammalian species including rabbits, guinea pigs, rats and mice as well as humans following neurosurgical resections. Rats have been the most commonly used species with multiple different strains being employed. However, the relative ease of modifying the mouse genome and the recent widespread use of transgenic mice strains has increased the popularity of mouse models for experimental use. The comparatively small size of the mouse brain implies that a single slice from a mouse brain is likely to preserve more

functional connectivity than a slice of similar thickness from larger mammalian species. Indeed, it is easier to induce seizure-like activity in hippocampal slices from mice than those from rats or humans⁵. However, the choice of species will depend on the specific scientific question asked.

Strains—Mouse strains display different sensitivity to epileptogenic conditions^{6,7} or may carry developmental deficits or characteristics that result from the different genetic background. The same is true for rats and other rodent species.

Sex—The background excitability of *in vitro* brain slices varies in male or female animals according to their age and hormonal state. Sexual differentiation is evident from very early in life and can be due to genetic factors, organizing effects of sex hormones (e.g., pre- or perinatal testosterone surge) and epigenetic factors. Additionally, sex differences in various signaling pathways controlling neuronal and glial activity and function, cellular morphology, synaptic connectivity and structure and function of various brain regions are known to exist⁸. Particularly (but not exclusively) in the context of mature animals, sex hormones, such as estrogen, progesterone or testosterone, modulate network excitability by interacting with sex hormone receptors, which are widely distributed throughout the brain and may affect seizure susceptibility^{9,10}. The most obvious evidence that sex-specific factors influence seizure susceptibility is catamenial epilepsy, which is present in approximately 40% of women with epilepsy¹¹. In this condition, seizure frequency varies during the ovarian cycle. Data from animal studies has demonstrated that the sex of the animal, and the levels of sex hormones can influence seizure frequency and induction although the precise direction of effects is varied and complex¹². Sex differences have also been described in *in vitro* recordings from immature animals¹³.

Age—The emergence of epileptiform activity *in vitro* is strongly affected by the age of the animal from which brain tissue is prepared^{14,15}. Nonetheless, the relationship between the age of animals and the ease of inducing seizure-like events *in vitro* is not a simple one. For example, in rat hippocampal slices, using the high K⁺ model, seizure-like events are not evoked in tissue from animals younger than postnatal day 5 (P5). These are easiest to evoke at P12 before becoming difficult to generate in tissue from animals over P21¹⁴. Such complexity emerges because age-related differences in excitability and seizure propensity are governed by multiple mechanisms¹⁵. As a single example, in contrast to adult tissue, GABAergic signaling is thought to be depolarizing early in development due to high levels of intracellular chloride¹⁶. It is important to note that the maturational trajectories and changes of various signaling pathways occur at different rates across species, strains or sex¹³. Rat and mice pups at P7–P12 are considered equivalent to human babies at birth. This is based on crude measures of brain growth rates, DNA, cholesterol or water content¹³. Furthermore, pubertal changes start around P32–36 in female rats and P35–P45 in males. It is therefore important that these trajectories, and the factors that control them, are considered when age groups are defined, that experimental groups are properly randomized for the ages of animals, and that the effect of age is adequately incorporated into study statistics and reported in the published manuscripts.

Animal breeding, housing and transport

Within a given strain, experimental results may vary depending on the breeders, or even on the geographical location of the same breeder¹⁷. How animals are bred, transported to and housed in the laboratory prior to the preparation of *in vitro* seizure models is seldom described even though these may have a considerable impact on cellular and network phenomena affecting epileptogenesis or ictogenesis. These factors include foster versus biological parenting, degrees of socialization, lack of environmental enrichment, or undue stress precipitated by poor climate control, maternal separation, handling or transport. For example, early life maternal separation and handling of neonatal pups has been shown to affect GABA_AR responses and cation chloride co-transporter expression in CA1 pyramidal neurons¹⁸, as well as to facilitate epileptogenesis following kindling¹⁹. Furthermore, environmental enrichment has been shown to be protective against kainate induced seizures²⁰. Regulation of lighting and climate is also important to standardize across experiments and optimize for the wellbeing of the animals. Rats are nocturnal animals and exposure to light for variable periods before euthanasia may generate different levels of stress. Similarly, exposure of rodents to temperatures above their thermoneutral zone (ranges between 26–30°C in rats; varies with activity level), which can happen in laboratories without adequate climate control, may also result in stress as rodents are unable to sweat and must resort to hyperventilation²¹. Neonatal pups, however, require raised ambient temperatures when separated from their litter, since nesting temperatures are higher.

Timing of brain tissue collection

Circadian rhythms and sleep wake cycles are important for network activity and synaptic plasticity, although the precise effects and their direction vary across species and brain areas²². Clinical and experimental research has demonstrated a clear relationship between sleep or wakefulness and seizure threshold. For example, in a rat model of temporal lobe epilepsy, seizures occurred more frequently during the light part of a 12 hour light/dark cycle²³. To our knowledge, it has not yet been conclusively demonstrated that the time an animal is killed relative to the light/dark cycle influences the propensity to elicit seizure-like events *in vitro*. However, various neurotransmitter systems are known to be modulated by the light-dark cycle, for example adenosine, an inhibitory neuromodulator has been shown to be more abundant at the start of the light cycle in rats²⁴.

In chronic animal models of epilepsy, spontaneous seizures may transiently change the excitability of neuronal networks and thus potentially influence research findings. For example, postictal refractoriness is associated with the prolonged activation of the adrenergic and GABAergic systems^{25,26}. Since the brain is never in a constant state, if only because of the circadian rhythm, there is a never a “baseline condition”. In animals with epilepsy, seizures induce another layer of complexity to the ongoing brain dynamics.

We suggest that investigators report detailed descriptions of how animals are bred, how long after transport animals were sacrificed, the type of and conditions of housing, the presence of environmental enrichment, light/dark cycle conditions and time of sacrifice of experimental animals relative to their light/ dark cycle, or other factors which may result in stress (see Figure 1).

We recommend reporting if and how seizure monitoring was done and the type of seizures captured prior to the tissue collection, to be aware of the time of the last seizure recorded in the animal before performing *in vitro* experiments. The time after a known seizure should be considered in the experimental design and data analysis.

Method of sacrifice and use of anesthesia

The use of anesthesia prior to decapitation is strongly advocated and often mandated by institutional or other governing bodies that ensure the humane handling of animals in experimental studies. For ethical reasons, and to diminish stress and pain, decapitation should be performed under anesthesia according to regulatory guidelines²⁷. It falls upon the investigators to justify and obtain approval for the exemption from using anesthesia prior to decapitation, when deemed inappropriate for the experimental goals. Direct application of anesthetics affects synaptic transmission with potential effects on seizure dynamics and electrophysiological responses of recorded cells²⁸. It is important to note that despite the effect of anesthesia on synaptic transmission, we do not actually know the extent to which anesthetics used at the time of decapitation might affect *in vitro* experiments performed remote to the decapitation. Residual effects have been documented in a study showing that capsaicin affected long-term potentiation in the lateral amygdala differently depending on the anesthesia (ether or isoflurane) used prior to decapitation²⁹. In contrast, comparison of the effects of various anesthesia protocols followed by decapitation on the electrophysiology of ischemic neocortical rat brain slices did not detect any differences among anesthetics³⁰. However, this finding may not exclude potential effects in other experimental settings assessing other markers of neuronal activity in naïve or non-injured tissue.

The most commonly used anesthetic strategies consist of intraperitoneal injection of barbiturates (pentobarbital) or the inhalation of volatile anesthetic agents (halothane, isoflurane, sevoflurane, or desflurane, with or without N₂O₂). Injectable barbiturates act quickly and reliably to render rodents unconscious. However, restraint is necessary and pain may be associated with injections given via the intraperitoneal route. Furthermore barbiturate anesthesia has the limitation that females often require higher doses to achieve adequately deep levels of anaesthesia⁵.

Therefore, the use of volatile agents, such as isoflurane, combined with a vaporizer is the preferred method of anesthesia. Restraint of the animal is not required and this method allows precise control of the depth of anesthesia before decapitation. In addition, the active concentration of volatile anesthetics in resected brain tissue is likely to decline more rapidly than other agents due to their low solubility and high volatility. The mildest form of anesthesia is probably isoflurane combined with nitrous oxide and 30% oxygen. Prolonged deep isoflurane anesthesia can however cause opening of the blood-brain barrier (BBB)³¹.

It is important that whatever method is chosen, the level of anesthesia is monitored and an appropriately deep level of anesthesia is reached before the animal is decapitated. The animal's muscles should be relaxed, withdrawal reflexes should be absent (ear pinch, toe pinch) and respiration should be within the normal range or slightly decreased (normal 70-115 breaths/min). Too light a level of anesthesia before performing decapitation could result

in undue pain and distress in the animal. In contrast, anesthesia that is too deep may result in circulatory arrest, death and a compromise of brain-tissue viability.

For older animals, in an attempt to enhance tissue viability, some investigators perform transcardial perfusion with a cold solution containing low calcium and reduced sodium prior to decapitation. Cardiac perfusion is thought to improve the quality of the slice, as assessed by ability to perform dendritic recordings and can be especially useful in adult or aged animals.

It is worth appreciating that when experiments are performed on resected human brain tissue from patients who have undergone epilepsy surgery, the patient has typically received a cocktail of anticonvulsants and anesthetics. We suggest waiting a minimum of 3 hours before eliciting reliable seizure-like events from human brain slices. In addition, the patient's sex and anticonvulsant use prior to surgery should be reported.

We suggest that anesthesia (type, route, dose) prior to decapitation should be well documented and standardized across experiments to optimize comparisons of *in vitro* electrophysiology studies. The selection of or exemption from anesthetic use prior to decapitation should be carefully determined and justified by prior literature or studies, the experimental goals, and in compliance with the regulatory guidelines (see Figure 1).

Brain slice preparation

All aspects of brain slice preparation influence the viability of brain slices. Optimal techniques often depend on the species and age of the animal or the brain area selected. Furthermore, strategies to improve slice health are under continual refinement and debate between laboratories. An exhaustive analysis into the possible drawbacks or benefits of specific manipulations is beyond the scope of this article. We, however, describe recent trends and explore issues that may be of possible relevance to those wishing to utilize *in vitro* models of seizures.

Slicing techniques and composition of slicing and recovery solutions—Early studies utilized the same artificial cerebral spinal fluid (aCSF) solution for cutting and storing slices as was used for recording purposes. Over the past two decades, various protective cutting methods have been developed for preparing healthy brain slices from juvenile and adolescent animals^{32,33}. These methods are based on the idea that passive sodium and chloride influx and subsequent cell swelling during slice cutting and recovery is the predominant insult that leads to reduced survival of neurons. This issue is especially important for those cells close to the slice surface, which are most likely to have sustained injury during the slicing process. The most commonly used protective technique first developed over two decades ago is to replace sodium chloride with equiosmotic concentrations of sucrose³². Variations of the standard sucrose protective slicing method have since been described. These include modified sucrose cutting aCSF regimes with optimized osmolarity³⁴, mixed NaCl/sucrose³⁵ or substitution of sodium using choline³⁶ - although it should be noted that at high concentrations (> 2mM) choline activates nicotinic and muscarinic receptors³⁷, N-methyl-D-glucamine (NMDG)³⁸, glycerol³⁹, or K-Gluconate⁴⁰. Recently, an optimized method employing NMDG as a sodium substitute

during both the cutting and recovery period has gained widespread popularity for preparing viable slices and acquiring intracellular recordings in brain tissue from mature and aged rodents^{33,41}. Maintaining cell viability in such tissue was previously exceptionally challenging.

Recent improvements in vibratome design and performance have also increased the quality of brain slice preparations. In particular vibratomes with minimal z-axis deflection are able to reduce damage to more superficial cells and structures³³. Following the cutting of brain slices, tissue is stored within either standard recording aCSF or specialized recovery aCSF (see above) inside either interface or submerged recovery chambers to enable the slices to 'recover' before recording. Likewise, various laboratories utilize different regimens in terms of length of recovery period and temperature at which slices are allowed to recover.

Nonetheless, it is clear that the combination of protective cutting solutions and enhanced vibratome performance have improved success rates for establishing recordings from delicate structures such as axons and dendrites^{35,42}. What is less certain is whether these advances in brain slice preparation play a role in the context of *in vitro* models of seizures. Indeed, there is currently considerable uncertainty as to whether improved slice viability enhances or reduces the likelihood of evoking epileptiform activity *in vitro*, particularly in tissue from naive animals. This stems from the inherent limitations of the acute brain slice preparation. Firstly, the tissue has experienced a period of ischemia. Secondly, projection fibers entering and leaving the slice have been severed. Thirdly, cells themselves are likely to have gone through varying degrees of physical or osmotic trauma during the slicing procedure. These processes are unlikely to affect excitatory and inhibitory systems to an equal extent. For example, in most brain regions, particularly the neocortex, the majority of inhibitory circuitry is local, whilst excitatory projection fibers between areas provide a major source of synaptic excitatory drive⁴³. These fibers are largely severed during slice preparation, which explains the drastic reduction in spontaneous synaptic activity as compared to the situation *in vivo*. Indeed, in all acute slice preparations, ictal events do not occur spontaneously but require the addition of pro-ictogenic agents to either enhance excitation or reduce inhibitory systems. Consistent with this, the threshold for inducing seizure-like events differs between preparations that include different brain areas. For example, transverse slice preparations which include the entorhinal cortex and hippocampus, thereby maintaining a large degree of intra and inter-area connectivity, are more amenable to induction of epileptiform activity⁴⁴.

Intriguingly it appears that inhibitory circuitry is more susceptible to damage during the slicing procedure than excitatory transmission. For example, Tanaka et al.,³⁸ demonstrated that the use of protective cutting solutions greatly enhanced the viability and survival of GABAergic interneurons in cortical slices from adult mice, in addition to promoting the health of primary glutamatergic neurons. In addition, Kuenzi et al.,⁴⁵ showed that long-term potentiation is reduced in hippocampal slices prepared using sucrose due to enhanced maintenance of inhibitory circuitry. It appears that the ability of fast-spiking interneurons to maintain rapid firing rates is also particularly sensitive to the reduced oxygen tension often present in acute slices. Furthermore, chloride influx is an inevitable consequence of the cellular damage that occurs during brain slice preparation, particularly within superficial

layers⁴⁶. This widespread intracellular chloride accumulation, results in a depolarizing and even excitatory shift in the effect of fast GABAergic transmission. While a certain level of brain slice health is required to evoke epileptiform activity *in vitro*, somewhat counter-intuitively for the reasons described above, developments which better preserve slice viability may ultimately reduce the likelihood of generating *in vitro* seizure-like events by protecting endogenous inhibitory function. Nonetheless, this is important as seizure-like events elicited within relatively intact inhibitory circuitry will better represent the *in vivo* and clinical situation.

A final consideration is the thickness at which brain slices should be prepared. When considering an optimal thickness there is a necessary tradeoff between preserving connectivity and maintaining adequate oxygenation in all areas of the slice. Thinner slices will be better oxygenated with poorer connectivity whilst thicker slices will preserve greater connectivity, but central areas are more susceptible to hypoxia. The use of interface or dual perfusion chambers with enhanced tissue oxygenation capability allow for thicker slices to be utilized. In general, investigators use slices with a thickness of between 350 and 450 μm .

We recommend that investigators report the type of chamber, as well as time periods and temperature used for recovery as these may affect various properties of brain slice function. For example, aspects of long-term potentiation induction in rat hippocampal slices may differ depending on whether slices recovered in an interface as compared to submerged environment⁴⁷.

We recommend that investigators consider using protective cutting solutions in order to better preserve brain circuitry following the slicing procedure. This is particularly relevant in preparations made from adult and aged rodents as well as surgically resected human tissue. Consideration and reporting of factors influencing viability and quality of slice recordings (e.g., thickness, perfusion speed, temperature, above and below slice perfusion and fluid level, pH) is strongly encouraged.

Organotypic slice cultures—Organotypic slice cultures represent an acute slice whereby the recovery period has been extended to span days, weeks and even months by maintaining the slices in an incubator with access to a culture medium. In many respects, acute slices represent a dynamic system whereby a subset of cells is gradually dying whilst the remaining neurons are recovering from the trauma associated with the slice procedure itself. Within an organotypic slice culture however, at least at the time of recording, cell viability has reached a relatively stable level. Organotypic slice cultures have been made from almost all brain regions including, hippocampus, cortex, cerebellum and brainstem structures. A drawback of this technique is that very young animals (P0 – P10) need to be used. Whilst the loss of specific cell types have not been observed following the culturing process⁴⁸ and the properties of synaptic transmission are generally maintained⁴⁹, considerable synaptic rearrangement does occur during the regrowth that follows slicing-induced deafferentation. For example, mossy fiber sprouting has been demonstrated to occur within organotypic hippocampal cultures⁵⁰. In general, recurrent connectivity increases as a function of time in culture, which is thought to underlie the gradual development of epileptiform activity in these preparations. Indeed, interictal-like population spikes develop

over a period of roughly two weeks which is then followed by the generation of spontaneous seizure-like events⁵¹. This stereotypical progression makes this preparation a potentially useful model for investigating the mechanisms underlying epileptogenesis – with particular significance as a model of post-traumatic epilepsy. An additional important advantage of organotypic slices is that the lengthy periods for which slices can be maintained allow for prolonged experimental access *in vitro*.

Although multiple methods exist to generate organotypic slice cultures, the simplest and most popular is the interface method first described by Stoppini et al.,⁵² and described in detail by De Simoni and Yu⁵³. This protocol however omits the neuronal culture supplement B27 (Gibco) which appears to be important for supporting the neuronal growth and viability required for the generation of spontaneous seizure-like events using these cultures. In addition, this protocol includes the use of antibiotics, which may affect neuronal activity and glial function. This points to an important disadvantage of this preparation: organotypic slice cultures are very sensitive to the multiple possible variations in culture conditions and the number of days spent *in vitro*. For example, difference between batches of horse serum or B27 may have considerable effects on the quality and levels of activity generated within cultures. Finally, investigators should be aware that organotypic slice perfusion during interface chamber recordings with 95% O₂/5% CO₂ (as opposed to air) has been associated with oxygen toxicity and reduced slice viability⁵⁴.

Optimization and reporting of highly detailed protocols for the generation and use of organotypic slice cultures is recommended to enhance reproducibility when using a preparation with considerable intrinsic variability. Reporting of the composition of culture media (e.g., growth factors, hormones, antibiotics) is important given their potential effects on recordings.

Intact preparations

In vitro preparations that preserve connectivity between distant areas of the brain were developed in the late 1980s to extend the study of intracerebral networks beyond the limitations of *in vitro* slice preparation. To maintain the oxygenation of large portions of the brain *in vitro*, either superfusion of embryonic *in toto* preparations (such as the whole hippocampus), or perfusion via the preserved vascular system in adult animals (as in the isolated guinea pig brain) was achieved. The specific methodological issues associated with the use of these two preparations will be reviewed in the next paragraphs.

Whole hippocampus—The whole hippocampus and related ‘intact structures’ can be investigated *in vitro*^{2,55,56}. These preparations are viable for electrophysiological and imaging experiments. Their major advantage over slices is that intrinsic connectivity is preserved. This maintenance of complex networks is useful for investigating seizure propagation, as seizure initiation and propagation zones can be manipulated independently⁵⁷. Another advantage of the intact preparation is the preservation of cell integrity. In slices, cells close to the surface swell and accumulate chloride, which does not happen in intact structures⁴⁶.

The main limitation is oxygen penetration, which drops considerably with depth⁵⁸. To solve this issue it is useful to use a double perfusion chamber, which provides oxygenation to both bottom and top parts of the preparation^{59,60}, as well as to employ fast perfusion speeds (10 ml/min). Since all it takes is some surgical skill to extract intact structures, complex networks can be studied, like inter-hemispheric⁶¹ or septum-hippocampus⁵⁷ communication. We find that tissue from more mature animals degrades more quickly, hence tissue from juvenile animals is typically used. P14 animals can be reliably recorded and imaged for up to 6 hours⁶². Although blind patch clamp recordings can be performed, the thickness of the tissue does not allow targeted recordings below the principal cell layer in the hippocampus. Nonetheless, 2-photon imaging can be performed at considerable depth⁶².

We recommend the use of this preparation when intact intra-hippocampal connectivity is required.

Whole brain—The isolated brain of the adult guinea pig maintained *in vitro* by arterial perfusion³ complements the obvious advantages of *in vitro* preparations (mechanical stability, easy access to brain tissue and pharmacological manipulations) with the complete preservation of the neuronal connectivity between distant brain regions associated with the functional integrity of the BBB. The preparation is viable for neurophysiological and imaging studies, in particular in ventral surface areas that are difficult to reach *in vivo*, such as the olfactory and limbic cortices. The access to deep brain regions is based on the use of external reference points and on the guide of stimulus-evoked field responses. The guinea pig brain can be isolated *in vitro* because of the peculiar arrangement of the communication between the vertebro-basilar and the carotid arterial systems that form the Willis circle that allows the perfusion of the entire brain via the basilar artery. In both rat and mouse, the small diameter of the posterior communicating arteries does not allow good brain perfusion *in vitro* when the basilar artery is cannulated. The facilitated access to both neuronal and vascular compartments in this preparation is ideal to induce acute changes that mimic neurological disorders, such as seizures, acute ischemia, brain inflammation, and is suitable for studies on the role of BBB in pathophysiology of brain diseases and for screening studies on BBB permeability of new drugs and neuroactive compounds.

Barbiturate anesthesia is utilized for surgery. The anesthetic is efficiently and completely washed out of the preparation after 1 hour *in vitro*⁶³. The method for brain extraction is not very different from the standard procedure utilized for *in vitro* slice preparation. However, a plasma expander (3% dextran 70,000) is added to the saline solution to maintain the osmotic properties of the intravascular compartment. Intracardiac perfusion with cold (10°C), carboxygenated solution precedes surgery, to reduce brain temperature and metabolism. After isolation and transfer in the incubation chamber the brain is perfused through the resident vascular system with solution at 7 ml/min. The temperature of the isolated brain is slowly increased to 32°C within one hour to coincide with washout of the anesthetic.

We recommend the use of this preparation to study interactions between remote brain regions and to simultaneously analyze neuronal and vascular compartments.

Recording conditions

Composition of recording solutions

Ionic composition—The composition of the aCSF utilized during the acquisition of experimental data varies greatly between laboratories. Even small differences in the aCSF ionic composition can have marked effects on the levels of excitability and network activity detected in any particular preparation. Indeed, many *in vitro* models of ictogenesis deliberately modify the composition of aCSF in order to elicit seizure-like events in otherwise normal tissue.

A ‘physiological’ aCSF is typically composed of the following: the predominant ions Na^+ and Cl^- at a concentration of between 120–140mM, 3–3.5mM K^+ , between 1.2–1.3 mM PO_4^- , 10–25 mM glucose, 22–26 mM HCO_3^- , 1–2 mM Mg^{2+} , and 1–3 mM Ca^{2+} . Particular care should be taken when adjusting the concentration of K^+ and the cations Ca^{2+} and Mg^{2+} , because even very small variations in the levels of these ions results in significant effects on neuronal activity. We recommend a K^+ concentration of 3 mM, although the actual concentration may be slightly higher (3.3 – 3.5 mM) *in vivo* in the awake behaving animal⁶⁴. aCSF concentrations of K^+ above 5 mM are commonly used to generate *in vitro* epileptiform activity. For a physiological aCSF, we advise the use of a Mg^{2+} concentration of 1.6 mM although magnesium concentrations of between 0.9 and 1.2 mM have been recorded from cerebrospinal fluid samples⁵. Reducing the Mg^{2+} concentration has a facilitating effect on synaptic transmission and below 0.9 mM can elicit epileptiform activity. Indeed, nominally zero Mg^{2+} aCSF is one of the most popular methods for generating *in vitro* seizure-like events. These events are thought to occur via a reduction in surface charge screening, and the Mg^{2+} dependent block of NMDA receptors at hyperpolarized potentials⁶⁵. The concentration of calcium *in vivo* is thought to be approximately 1.2 mM, which is mimicked by an *in vitro* concentration of 1.6 mM as HCO_3^- chelates about 25% of the free calcium⁵. Varying the extracellular calcium concentration has a complex effect on synaptic transmission and neuronal activity. For example, utilizing calcium free media is another popular method of evoking epileptiform activity. Despite the complete abolishment of synaptic transmission, the epileptiform activity is thought to occur via reduced surface charged screening and Ca^{2+} sensitive potassium currents combined with enhanced synchrony via ephaptic coupling⁵. In contrast, aCSF with raised calcium can also enhance neuronal activity by promoting long term potentiation⁶⁶.

Amino acids are rarely added to the aCSF although they are typically found in appreciable concentrations within the interstitial space of the nervous system. Examples include glutamine (0.5 mM) and GABA (20 μM). Furthermore, it is currently open to debate as to whether young tissue might require additional energy substrates, such as ketone bodies^{67,68}.

It is important to note that the 10–25 mM concentration of glucose used in aCSF is not physiological, as average glucose concentrations vary (according to the technique used) between 0.8–2.3 mM in the brain^{69,70}. The use of 10–25 mM glucose effectively clamps the delivery of energy substrates.

Osmolarity—It is important that the osmolarity of the recording solution is adjusted carefully as changes in osmolarity can have profound effects on network excitability. A typical aCSF osmolarity should approach 290 mOsm. Reductions in the osmolarity (e.g. a 35 mOsm reduction) of recording solutions causes cell swelling, shrinkage of the extracellular space, enhanced excitability and the emergence of epileptiform activity presumably by intensifying ephaptic interactions⁷¹. Raised osmolarity has the opposite effect, with consequent cell shrinkage having an anticonvulsant effect.

pH—The negative logarithm of H⁺ ion concentration (pH) is a fundamental parameter with powerful effects on synaptic transmission and network excitability. It is well known that more acidic recording solutions reduce excitability, whilst more alkaline solutions promote hyperexcitability⁷². These effects are thought to be mediated by a variety of processes. Changes in pH have been shown to alter the conductance of many neurotransmitter receptors. For example, acidic and alkaline shifts have been shown to reduce and enhance the permeability of NMDA receptors, respectively⁷³; whilst opposite pH-induced changes to conductance have been demonstrated for GABA_A receptors⁷⁴. Acidic shifts have also been shown to enhance the release of adenosine with concomitant reduction in network excitability⁷⁵. Investigators should attempt to maintain the pH of recording solutions within 0.2 pH units of 7.4. This is typically achieved by bubbling a bicarbonate buffered recording solution with carbogen (95% O₂/5% CO₂). It is important to note that CO₂ dissolves more readily in cold as opposed to warm solutions. As dissolved CO₂ results in the production of carbonic acid, this temperature dependence means that changes in aCSF temperature affect the pH of experimental solutions. We find that when experimental solutions are maintained at 35 degrees, a concentration of 21 mM NaHCO₃ results in a pH of 7.4. When working at room temperature, a NaHCO₃ concentration of 26 mM is more appropriate for maintaining extracellular pH at 7.4. Investigators should be aware that the use of HEPES buffered solutions at a pH of 7.4 (instead of NaHCO₃) reduces slice excitability via intracellular acidification of neurons⁷⁶.

Temperature—Temperature is a potent modulator of neuronal activity. In addition to affecting pH (see above), the primary manner in which temperature affects network function is by altering the reaction kinetics of a wide array of proteins and channels. For example, the kinetics of voltage gated sodium and potassium channels are altered by changes in temperature with profound consequences for spiking activity⁷⁷. As a result, the temperature of the recording solution is important for controlling the inducibility of epileptiform activity when employing *in vitro* models of seizures. Below 32°C, seizure-like events are often difficult to elicit⁷⁸. In addition, rapid cooling is sufficient to terminate seizure-like events *in vitro*⁷⁹. Conversely raising the temperature of the experimental medium above 38.2°C is able to generate epileptiform activity in otherwise normal tissue⁸⁰. Note that the use of near physiological temperatures decreases the survival time of slices.

It is critical that investigators carefully monitor and report the ionic composition, osmolarity, pH and temperature of recording solutions as these parameters have marked effects on the emergence and nature of *in vitro* epileptiform activity (see Figure 1).

Recording chambers and perfusion speed

There are two main types of recording chambers utilized for *in vitro* research on brain slices; interface and submerged chambers. When using interface chambers, the brain slice is maintained on an interface between the recording solution below and humidified and oxygenated gas above. The advantage with this approach is that slices are well oxygenated and equally perfused. This means that relatively slow perfusion rates (1 – 2 ml/min) are sufficient to maintain slice health and elicit epileptiform activity when employing *in vitro* models of seizures. A significant disadvantage is that water immersion objectives that are necessary for making visually-targeted patch-clamp recordings cannot be utilized with interface chambers, limiting their use to the measurement of local field potentials and intracellular recordings with blind patching or sharp microelectrodes. For this reason, submerged chambers are considerably more popular. In this configuration the brain slice typically rests on the base of the chamber where it is submerged within the perfusate. It has now been relatively well documented that higher perfusion speeds are required for adequate tissue oxygenation under these conditions^{59,78}. It should be noted that important differences in various slice parameters such as ion homeostasis mechanisms and responses to anoxia exist between interface and submerged chamber recording conditions^{81,82}. Investigators should be aware of these when studying *in vitro* epileptiform activity using either type of chamber.

When using submersion chambers, we recommend using perfusion speeds of at least 4 – 5 ml/min in submerged slices in order to elicit epileptiform activity. Furthermore, we recommend minimizing the level of fluid above the slice in the submersion chambers as this tends to improve oxygenation and tissue viability. It is worth appreciating that the microscope objective inserted into the fluid above the slice may result in a zone of slowed perfusion. In these cases, the use of a dual perfusion chamber, which enables oxygenation of both bottom and top parts of the slice⁵⁹, should be strongly considered. Sophisticated triple chambers have also been developed which allow for recording from the intact hippocampal formation whilst allowing for the selective application of pharmacological agents to either hippocampus or connecting commissural fibers².

Investigators should remain cognizant of the fact that each preparation has an optimal time window during which reliable recordings can be made. Therefore, this should be considered and reported.

Electrophysiological recording methods

The grease gap chamber—The simplest device for recording of neural activity is the grease gap chamber. A brain slice is placed over a divide between two chambers, which are individually perfused. These two chambers are isolated by silicone or vaseline. Proconvulsant agents can be applied to one chamber. If this solution induces epileptiform activity, a current will flow between the chambers and the resulting potential difference can be recorded using electrodes inserted in each of the two chambers⁸³.

Glass pipette based recording techniques—Field potential recordings represent the cumulative electrical activity from the surrounding neuronal population. As such, this

technique is useful for recording the synchronized neuronal activity which constitutes epileptiform events. Glass pipette recordings for field potentials use electrode tips between 1 – 5 μm filled with physiological NaCl or artificial cerebrospinal fluid solution (aCSF). Electrical contact to an amplifier is made using a silver chloride wire. However, if these are oxidized they can become light sensitive. Glass electrodes of this nature allow recordings of single unit activity, field potentials and direct current coupled potentials to be made in either interface or submerged recording chambers. This can be combined with higher resolution techniques including whole-cell patch-clamp or sharp microelectrode recordings, which allow for the measurement of membrane potential and/ or synaptic currents. However, these techniques typically require visual guidance using submerged chambers and water immersion objectives for single-cell or subcellular targeting. It is worth noting that when performing slice recordings, cells nearest to the slice surface are likely to be damaged with high levels of intracellular chloride; recording cells at least 50 μm below the slice surface may therefore avoid traumatized neurons⁴⁶. A further advantage of submerged chamber recordings is that one can identify the cells from which recordings were made by filling them with dyes. In addition, specific cell-types may be targeted for recordings if transgenic mouse lines are utilized where the expression of fluorescent reporters is under the control of cell-type specific promoters. The development of the cre-lox system and the wide availability of multiple cre recombinase driver and cre reporter mouse lines have greatly facilitated the ability of investigators to generate targeted recordings from genetically defined cell types *in vitro*⁸⁴. It is important that investigators validate the cell expression profiles of cre mouse lines before use, as off target expression has been noted in some cases⁸⁵.

Reporting the type and characteristics of glass pipette electrode recordings is recommended. When using brain slices, recording from cells at least 50 μm below the slice surface is recommended to avoid traumatized neurons.

Carbon fiber and metal based electrodes—Recordings with carbon fiber or metal electrodes enable the recording of local field potentials in addition to single unit activity - the extracellular equivalent of action potentials. Carbon fiber electrodes are usually able to detect more cells than metal electrodes. Carbon fiber electrodes can also be covered with enzymes, which permit the detection of extracellular transmitters. Metal based electrodes can be engineered to have multiple recording sites. This enables the recording of multiple units from a single electrode. Multi electrode arrays have also been developed and are particularly suitable for *in vitro* use where brain tissue can be overlaid on the array allowing for close contact between neurons and electrode sites. These devices permit recordings from multiple single units over relatively broad regions of tissue to be made.

Ion selective microelectrodes—Seizure-like events are associated with considerable ion fluxes across cell membranes which can be detected by using appropriate ion sensitive microelectrodes. These are usually two barrel glass electrodes where one barrel records the extracellular field potential and the other, which is filled with the appropriate ion sensitive resin, records changes in the ion concentration of choice and the field potential. The difference between the two barrels then represents changes in ion concentration. For

measurements of glucose and oxygen, Clark-type electrodes can be used which exploit the change in resistance of a platinum wire at a defined voltage which confers substrate specificity. Amperometric or voltammetric methods are available. These techniques can also be used to detect neurotransmitters such as dopamine or norepinephrine.

Functional microscopy and optogenetics—The development of optical imaging techniques such as confocal and multi-photon microscopy in combination with ion-sensitive dyes and ion-sensitive genetically-encoded indicators of ion concentration have rapidly gained popularity for reporting both ion dynamics, and by proxy, neural activity in the nervous system. Optical reporters of Ca^{2+} , Na^+ , Cl^- and pH have all been successfully utilized to quantify concentration changes in these ions during *in vitro* epileptiform activity in varying contexts^{86–89}. Beyond ion concentration, sensors have also been developed to report an ever increasing array of metabolites and neurotransmitters^{90,91}. A major advantage of all genetically-encoded sensors is that they may be genetically targeted to specific cell-populations or subcellular locales. In addition to sensors, the relatively recent development of genetically-encoded, light activatable ion channels now allows for optical control of genetically defined subsets of cells⁹². These optogenetic techniques can be used to determine the contribution of different cell types to *in vitro* epileptiform activity⁹³. Finally, optogenetic silencing strategies have been used in order to try and control epileptiform *in vitro*⁹⁴. However, it is important to note that the use of opsins can result in significant changes to ion gradients within brain tissue^{95,96}.

Concluding Remarks

In vitro models have been extensively utilized to investigate the mechanisms underlying seizures and epileptogenesis. Advances in brain slicing and recording technology mean that *in vitro* methods will continue to grow in relevance. In the course of this review we hope to have explored the multitude of important methodological considerations, which are required to effectively pursue this valuable avenue of research.

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Key points

1. In vitro preparations are an important means for understanding mechanisms of epilepsy and epileptic seizures.
2. We critically review the numerous in vitro methods utilized in epilepsy research.
3. We call for the inclusion of detailed descriptions of techniques and often ignored parameters.
4. We explore how recent developments in brain slice preparation affect their use as models of epilepsy.

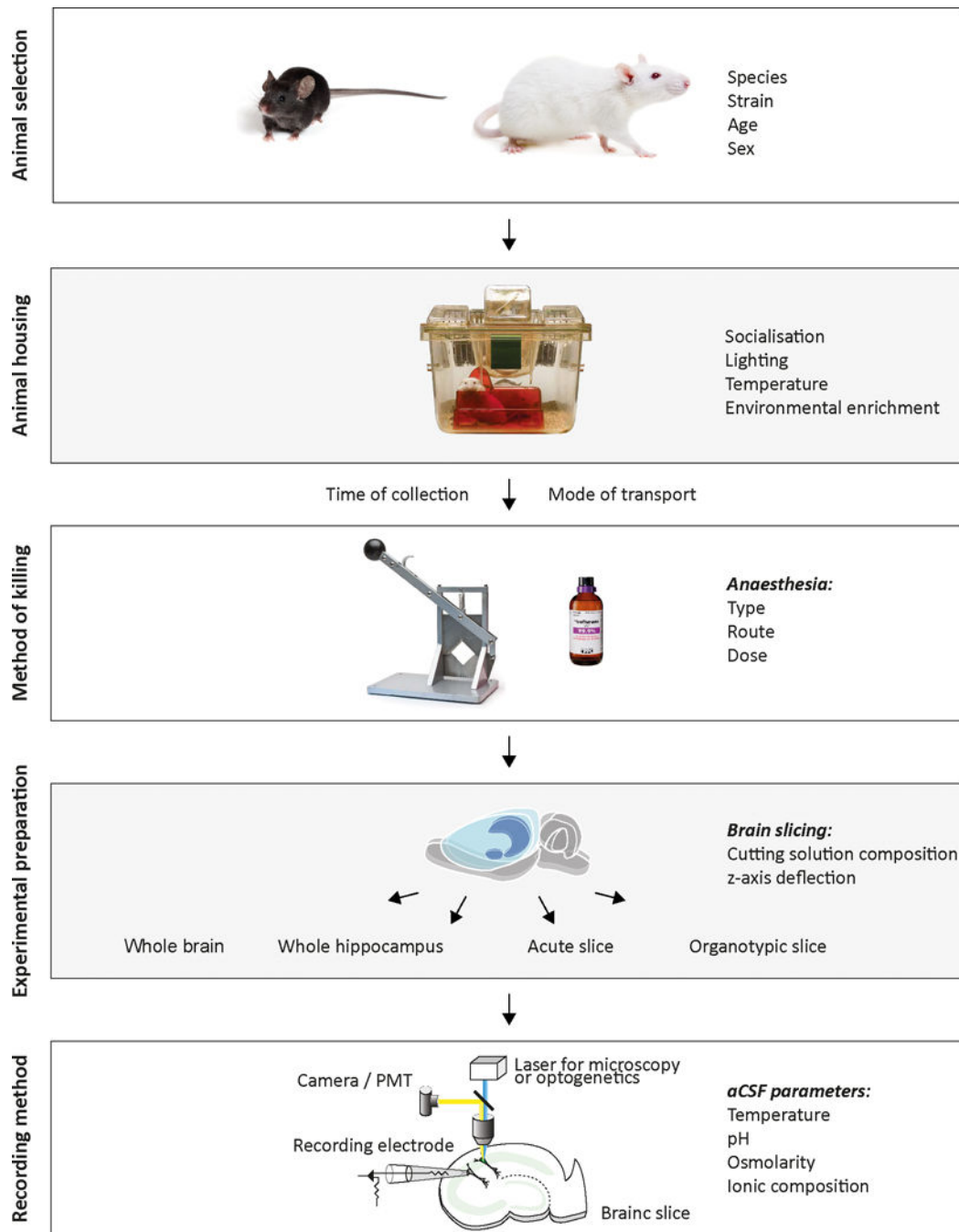


Figure 1.

Methodological parameters for *in vitro* models of seizures and epilepsy. The preparation of *in vitro* models of seizures and epilepsy involve multiple parameters, which may influence the validity and reliability of scientific findings. The species, age, strain and sex of experimental animals to be utilized should be carefully considered before beginning experiments. Animals should be appropriately housed taking careful note of socialization, lighting, temperature and environmental enrichment. The time of collection and mode of transport of animals prior to experimentation are important variables, which should be

reported where possible. We suggest that the type, route and dose of anesthesia prior to decapitation be well documented and standardized across experiments. The selection of or exemption from anesthetic use prior to decapitation should be carefully determined and justified by prior literature, the experimental goals, and be in compliance with regulatory and ethical guidelines. Multiple different preparations may be used for *in vitro* studies of seizures and epilepsy including whole brain, whole hippocampus, acute and organotypic brain slices. When preparing brain slices, investigators should consider optimizing cutting solutions and the z-axis deflection of vibratomes in order to maximize slice health. Multiple techniques exist for monitoring and manipulating *in vitro* epileptiform activity. Investigators should carefully monitor and report the temperature, pH, osmolarity and ionic composition of recording solutions as these variables have marked effects on the emergence and nature of *in vitro* epileptiform activity.