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1. Summary of all recommendations for IMPROVE (Ischaemia Models: Procedural Refinements Of In Vivo Experiments)

The recommendations describe opportunities to improve the in vivo modelling of ischaemic stroke and minimise the level of severity in the most common rodent models of cerebral ischaemia, while sustaining or improving the scientific outcomes. The aim is to provide support for researchers and animal care staff to refine their procedures and practices, and implement small incremental changes to improve the welfare of the animals used and to answer the scientific question under investigation.

TIMING		RECOMMENDATION	TIMING		RECOMMENDATION		
Basic requirements before stroke	1	Rodents ordered from an outside supplier should be delivered at least seven days before the procedure	Intraoperative care	24	Aseptic surgical technique is essential.		
		to allow acclimatisation to the new environment.		25	Antibiotics should not be used prophylactically ur		
surgery	2	Animals should be acclimatised in harmonious groups before the start of the experiment. Re-housing animals in new groups should be avoided.		26	The surgeon should work with an assistant.		
	3	Animals should be acclimatised to handling and should not be handled by the tail. Tunnel and cup		27	Surgeons' performance should be monitored and		
		handling should be used for mice; rats should be handled by grasping around the shoulders.		28	During general anaesthesia and in the immediate p should be maintained by insulation or supported b that cuts out when normal body temperature is rea		
	4	Animals should be weighed daily for at least three days before surgery.	Post-operative care				
	5	Cage substrate, nesting material and shelter are basic welfare needs for rodents and should be provided. Tunnels, wheels and chewing sticks are simple, cost-effective ways to improve enrichment.		29 30	Additional care and monitoring of body temperatu		
	6	Additional 'super-enrichment' should be considered carefully, as it may have neurorestorative effects. Enrichment should be reported in publications specifically.			Monitoring of respiratory and cardiovascular para reproducibility of study methods.		
	7	In consultation with the veterinary and animal care staff, consideration should be given to the bedding materials and any new material should be introduced prior to surgery to acclimatise the animals.		31	Minimum parameters to monitor are depth of anal activities should also be recorded.		
	8	After stroke, animals should be returned to the same group of animals they were with before surgery as soon as they are sufficiently recovered.		32 33 34	Pulse oximetry is recommended as SaO ₂ is a goo difficult to monitor otherwise.		
	9	The randomisation protocol should ensure that each cage contains sham-operated and stroke animals, and/or animals allocated to different treatments.			Invasive monitoring is useful for experiments carr specific cases justified by the study needs.		
	10	Animals should have access to their post-stroke diet prior to surgery and surgery should not be undertaken until they reliably consume the diet.			Intubation and artificial respiration should be cons of ischaemic lesions, particularly for those lasting		
	11	Rodents should not be routinely fasted before surgery, unless there is a scientific reason; any restriction		35	Animals should be administered fluids pre-emptive		
	12	should be reported specifically in publications. After any food restriction for training purposes, sufficient time should be left to re-establish normal		36	The experimental study plan should include detail assessment points. This should be devised in con		
		feeding patterns before surgery.		37	All animals should be monitored frequently (at lea		
	13	Consistency of inter-animal housing, feeding and handling practices before and after stroke should be ensured.			post-stroke) using a traffic light system (see table pre-defined humane endpoint (red status).		
	14	Aged animals and those with co-morbidities should receive extra monitoring.		38	Monitoring frequency must be increased if co-mo clinical signs requiring intervention (amber status		
	15	Teeth should be checked regularly, especially if the animal is on soft food diet. Animals should be provided with chew sticks to grind teeth.		39	Monitoring duration should be long enough to ens		
Anaesthesia and analgesia	16	The anaesthetic should be chosen on the basis of both welfare and scientific outcomes and should take account of species, strain and health status of the animals. Selection should involve the vet.		40	Clinical assessment sheets should be completed remain with animal cages to ensure record of obs		
	17	Sham-operated animals should receive exactly the same anaesthesia regimen for the same duration as the test group in order to control for effects of the anaesthetic on outcomes.		41	Dehydration should be assessed frequently and t drinking normally.		
	18	Local anaesthesia should be used prior to incision during surgery, particularly if other types of analgesia		42	Additional hydration should be provided at least f		
		are not being provided, and with knowledge of local anatomy to ensure that it is applied in the appropriate area.		43	Animals should be provided with an appropriate p loose pellets on the cage floor should be supplied		
	19	Pain is a variable which needs to be controlled. Pain relief must be used unless there are good scientific reasons not to, supported by solid, reproducible evidence.					
	20	The analgesic drug should be selected in consultation with the vet, based on the objective of the study, the specific stroke model and the type and timing of outcome measures.					
	21	The animal should be assessed for level of pain post-operatively, to ensure that the analgesic regime is effective and to minimise the risk of any unnecessary medication or side-effects.					
	22	All animals should either receive the same doses of analgesics to avoid pain relief being a confounder, or the experimental design and analysis should account for animals receiving different doses. This should be reported explicitly in publications.	(Ischaemia Models: Pi		Sert N, Alfieri A, Allan SM, <i>et al</i> . The IMPROVE Guidelines rocedural Refinements Of <i>in Vivo</i> Experiments). <i>J Cereb Blood</i> II: <u>10.1177/0271678X17709185</u>		
	23	Analgesia should be given by the most reliable and least stressful route. If there is doubt about oral consumption, analgesics should be given parenterally.					

unless there is a justified case.

nd reviewed.

e post-operative period, the animal's body temperature d by a heating device, with a feedback heating system reached.

ature may be needed for obese animals.

rameters is essential for animal safety and the

aesthesia, respiratory rate and temperature. Surgical

ood indicator of tissue oxygenation and heart rate is

arried out under terminal general anaesthesia or for

onsidered for experimental protocols involving induction ng longer than 30 minutes.

tively to prevent dehydration during surgery.

ails of planned post-operative intervention and onsultation with veterinary and animal care staff.

east 4 times a day at regular interval during the first 48h le overleaf) and should be humanely killed if they reach a

norbid animals are used and if animals are showing any us).

nsure eating and drinking behaviours are observed.

ed each time animals are monitored – such sheets should oservations and consistency of care.

I treated post-operatively until the animal is seen to be

t for the first two days post-surgery.

post-surgical diet (e.g. wet mash). Softened food and ed for at least seven days post-stroke.





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2. Signs to monitor after experimental stroke surgery in rodent models

Clinical signs that may present following induction of cerebral ischemia have been characterized as green, amber or red, as detailed below. The guidelines apply to adult rodents from standard, commonly used strains and may need adjustment under other circumstances (e.g. for aged, young or obese animals).

	GREEN	AMBER	RED
	Clinical signs similar to those seen in clinical stroke are expected to be present after a focal ischaemic insult as it results in brain injury and swelling. For some studies where scientific endpoints can be achieved by small infarcts, signs may be reduced in severity or absent. It is anticipated that the majority of animals will not display all of the clinical signs, and any clinical signs should start to show improvement by 48h post-stroke.	The presence of any of the signs in the amber column below requires an increase in the frequency of monitoring and appropriate intervention, such as consulting the vet or implementing some of the recommendations presented overleaf. If any of the clinical signs exceed the limits stated, the animal should automatically move to RED status.	The presence of any of immediate euthanasia of killing (in the UK, Sched
TIMING			
Anytime	 Sensorimotor deficits (e.g. forelimb weakness/hemiparesis, altered gait) Neglect of one side of the body, preference for one direction of movement Reduced food and water intake Abnormal behaviour upon handling – this may involve increased or decreased reaction to being handled compared to non-stroke animals 	 Intermittent circling behaviour Absence of urine * Absence of faeces * Surgical wound complication Reduced motility, beyond what would be expected based on the severity of the ischaemic insult 	 Presence of barrel Presence of tonic of Continuous laboure normal for the age/
First 24h post-stroke	Lethargy and reduced motility with large strokes	 Animal not eating some normal chow or food supplement (e.g. wet mash) 	 Animal not moving, recumbent position recovery from gene duration of anaesth
First 48h post-stroke	 Weight loss ** Piloerection/ staring coat Discharge from the eyes and nose 	 Progressive weight loss exceeding 10% of the animal's pre-stroke weight Audible respiratory noises (rasping, wheezing), usually intermittent and normally not associated with an increased respiratory effort *** 	Weight loss of 20% decision not to eutl able to move aroun independently ende
First week post-stroke	 Recovery of weight loss starting 4 to 5 days post stroke Disruption to nest building activity during first week post stroke – although this is strain dependent as some strains don't nest build 	 Weight loss of up to 20% beyond 48h post-stroke (despite all efforts to supplement fluid and diet) but animal is eating well, drinking and able to move around and explore its environment. No recovery of weight towards pre-stroke level by 4 days post-stroke Intermittent abnormal motor activity, suggestive of seizure, during first 72h only Piloerection beyond the first 48h Reduced grooming beyond the first 48h Secretions around the nose and eyes persisting beyond first 48h No evidence of a return to near normal eating and drinking behaviour by day 4 post-stroke 	 Weight loss exceed efforts to suppleme if animal is eating w its environment) mu animal welfare offic No recovery of weig A decision not to eu and exhibiting signs endorsed by the ve Intermittent abnorn beyond first 72h Intermittent abnorn for two consecutive 48h ***

* These would typically be observed in singly housed animals to assess if the animal is eating and drinking sufficiently. However, group-housing has been shown to aid recovery. In group-housed animals, observing eating/drinking habits, monitoring body weight and whether the animals defecate/urinate during handling can be used to assess whether each animal is eating and drinking sufficiently.

** Weight loss after stroke is common, both in humans and in rodent models, and can be explained principally by dehydration, impaired feeding, inactivity and paralysis. However, other factors such as neuroendocrine sympathetic activation, fever and inflammation also contribute to metabolic imbalance and an increased catabolic drive leads to tissue wasting of both fat and muscle, depleting energy stores and leading to functional decline. Rodent models typically demonstrate a dramatic weight loss after stroke surgery, which normally starts recovering after 4-5 days. The amount of weight lost during that period is tightly correlated with the size of the infarct.

*** Abnormal breathing which is beyond what is considered normal for the age/weight and characteristics of animal exposed to stroke. For example, obese animals may exhibit noisier rasping respiration than non-obese animals. Intermittent wheezing (amber sign) can be seen in the first 24 hours after MCAO and is normally associated with intubation and/or long or repeated anaesthesia, most likely due to accumulation of respiratory secretions and/or minor laryngeal trauma. It should resolve within 24-36 hours. Respiratory distress (red sign) may be a result of pulmonary oedema associated with the MCAO or severe laryngeal trauma at the time of intubation.

of the signs in the red column below requires a of the animal via an approved method of humane edule 1 or other licensed method).

l rolling

- clonic seizures
- red respiration which is beyond what is considered e/weight of animal exposed to stroke ***

g, unresponsive to stimulation, or in a lateral on after the normal period of time expected for neral anaesthesia, given the anaesthetic used, the sthesia and the severity of the ischaemic insult.

% despite all efforts to supplement fluid and diet. A uthanise (e.g. if animal is eating well, drinking and und and explore its environment) must be idorsed by the vet or animal welfare officer

eding 20% beyond 48h post-stroke despite all ment fluid and diet. A decision not to euthanise (e.g. well, drinking and able to move around and explore must be independently endorsed by the vet or ficer

eight towards pre-stroke level by 7 days post-stroke. euthanise (e.g. if the animal is eating well, drinking ns of normal behaviour) must be independently vet or animal welfare officer

rmal motor activity, suggestive of seizure, persisting

ormal and laboured breathing (e.g. rasping) observed ive observations, or persisting beyond the first



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